Europäisches Patentamt

European Patent Office

Office européen des brevets



(11) EP 0 887 412 B1

(12)

# **EUROPEAN PATENT SPECIFICATION**

(45) Date of publication and mention of the grant of the patent:

15.10.2003 Bulletin 2003/42

(51) Int CI.7: **C12N 15/40**, C12N 7/01, A61K 39/12, G01N 33/569

(21) Application number: 98201704.8

(22) Date of filing: 22.05.1998

(54) Recombinant birnavirus vaccine

Rekombinanter Birnavirusimpfstoff Vaccin recombinant du birnavirus

(84) Designated Contracting States:

AT BE CH DE DK ES FI FR GB GR IE IT LI LU MC

NL PT SE

(30) Priority: 26.05.1997 EP 97201599

- (43) Date of publication of application: 30.12.1998 Bulletin 1998/53
- (73) Proprietor: Akzo Nobel N.V. 6824 BM Arnhem (NL)
- (72) Inventors:
  - Mundt, E.
     18461 Millienhagen (DE)
  - Lütticken, H.D.
     5831 CE Boxmeer (NL)
  - van Loon, A.A.W.M.
     5836 BB Sambeek (NL)
- (74) Representative:
   Mestrom, Joannes Jozef Louis et al
   Intervet International B.V.,
   P.O. Box 31
   5830 AA Boxmeer (NL)
- (56) References cited: WO-A-95/26196
  - E.MUNDT AND V.N.VAKHARIA: "Synthetic transcripts of double-stranded Birnavirus genome are infectious" PROCEEDINGS OF THE NATIONAL ACADEMY OF SCIENCES OF USA., vol. 93, October 1996, WASHINGTON US, pages 11131-11136, XP002044624

E.MUNDT ET AL.: "Identification of a novel viral protein in infectious bursal disease virus-infected cells" JOURNAL OF GENERAL VIROLOGY, vol. 76, no. 2, February 1995,

READING GB, pages 437-443, XP002044625

- E. MUNDT ET AL.: "VP5 of Infectious Bursal Disease Virus is not essential for virus replication in cell culture" JOURNAL OF VIROLOGY, vol. 71, no. 7, July 1997, pages 5647-5651, XP002044626
- MURPHY ET AL: 'PRODUCTION AND LEVEL OF GENETIC STABILITY OF AN INFLUENZA A VIRUS TEMPERATURE-SENSITIVE MUTANT CONTAINING TWO GENES WITH TW MUTATIONS' INFECTION AND IMMUNITY vol. 37, no. 1, 1982, pages 235 - 242
- Mundt E: An Abstract circulated at the Annual Meeting of the German Virological Society 1997 (Hamburg, 10-13 March 1997)
- Mundt E: A poster presented at the Annual Meeting of the German Virological Society 1997 (Hamburg, 10-13 March 1997)
- Mundt I: An Abstract in the Annual Report 1996 of the German Federal Research Institute for virus diseases in animal, published April 1997

#### Remarks:

The file contains technical information submitted after the application was filed and not included in this specification

P 0 887 412 B1

Note: Within nine months from the publication of the mention of the grant of the European patent, any person may give notice to the European Patent Office of opposition to the European patent granted. Notice of opposition shall be filed in a written reasoned statement. It shall not be deemed to have been filed until the opposition fee has been paid. (Art. 99(1) European Patent Convention).

### Description

10

20

30

35

40

[0001] The present invention is concerned with a bimavirus mutant, a vaccine comprising this mutant, a method for determining birnavirus infection in an animal, as well as with a test kit for carrying out this method.

[0002] Infectious bursal disease virus (IBDV) and Infectious pancreatic necrosis virus (IPNV) are members of the Birnaviridae family. Viruses in this family have a very similar genomic organisation and a similar replication cycle. The genomes of these viruses consist of 2 segments (A and B) of double-stranded (ds) RNA. The larger segment A encodes a polyprotein which is cleaved by autoproteolysis to form mature viral proteins VP2, VP3 and VP4 (Hudson, P.J. et al, Nucleic Acids Res., 14, 5001-50012, 1986; Dobos P., Annual review of fish diseases 5, 25-54, 1995). VP2 and VP3 are the major structural proteins of the virion. VP2 is the major host-protective immunogen of birnaviruses, and contains the antigenic regions responsible for the induction of neutralising antibodies. The VP4 protein appears to be a virus-coded protease that is involved in the processing of a precursor polyprotein of the VP2, VP3 and VP4 proteins. The larger segment A possesses also a second open reading frame (ORF), preceding and partially overlapping the polyprotein gene. This second open reading frame encodes a protein VP5 of unknown function that is present in IBDV infected cells (Mundt, E. et al., J. Gen. Virol., 76, 437-443, 1995).

[0003] The smaller segment B encodes VP1, a 90 kDa multifunctional protein with polymerase and capping enzyme activities (Spies, U. et al., Virus Res., 8, 127-140, 1987 and Spies, U. et al., J. Gen. Virol., 71, 977-981, 1990; Duncan R. et al., Virology 181, 541-552, 1991).

[0004] For IBDV, two serotypes exist, serotype 1 and 2. The 2 serotypes may be differentiated by virus neutralisation (VN) tests. Furthermore, subtypes of serotype 1 have been isolated. These so-called "variant" viruses of serotype 1 can be identified by cross-neutralisation tests (Diseases of Poultry, 9th edition, 1991, Wolfe Publishing Ltd, ISBN 0 7234 1706 7, Chapter 28, P.D. Lukert and Y.M. Saif, 648-663), a panel of monoclonal antibodies (Snyder, D.B. et al., Arch. Virol., 127, 89-101. 1992.) or RT-PCR (Jackwood, D.J., Proceedings of the International symposium on infectious bursal disease and chicken infectious anaemia, Rauischholzhausen, Germany, 155-161, 1994). Some of these subtypes of serotype 1 of IBDV have been described in literature for example: Classical, Variant-E, GLS, RS593 and DS326 strains (Van Loon, et al. Proceedings of the International symposium on infectious bursal disease and chicken infectious anaemia, Rauischholzhausen, Germany, 179-187, 1994).

[0005] Infectious Bursal disease (IBD), also called Gumboro disease, is an acute, highly-contagious viral infection in chickens that has lymphoid tissue as its primary target with a selective tropism for cells of the bursa of Fabricius. The morbidity rate in susceptible flocks is high, with rapid weight loss and moderate mortality rates. Chicks that recover from the disease may have immune deficiencies because of the destruction of the bursa of Fabricius which is essential to the defence mechanism of the chicken. The IBD-virus causes severe immunosuppression in chickens younger than 3 weeks of age and induces bursal lesions in chicks up to 3 months old.

[0006] For many years the disease could be prevented by inducing high levels of antibodies in breeder flocks by the application of an inactivated vaccine, to chickens that had been primed with attenuated live IBDV vaccine. This has kept economic losses caused by IBD to a minimum. Maternal antibodies in chickens derived from vaccinated breeders prevents early infection with IBDV and diminishes problems associated with immunosuppression. In addition, attenuated live vaccines have also been used successfully in commercial chicken flocks after maternal antibodies had declined.

[0007] Recently, very virulent strains of IBDV have caused outbreaks of disease with high mortality in Europe. The current vaccination programs failed to protect chicks sufficiently. Vaccination failures were mainly due to the inability of live vaccines to infect the birds before challenge with virulent field virus.

[0008] Eradication of the disease by other preventative measures than vaccination has not been feasible, because the virus is widely spread and because with currently administered live attenuated or inactivated IBDV vaccines it is not possible to determine whether a specific animal is infected with an IBDV field virus or whether the animal was vaccinated with an IBDV vaccine. In order to be able to start an eradication control programme for IBDV it is highly desirable that the possibility exists to discriminate between animals vaccinated with an IBDV vaccine and those infected with a field virus so as to be able to take appropriate measures, i.e. remove infected flocks, to reduce spreading of the virulent field virus. The introduction of, for example, a serologically identifiable marker can be achieved by introducing a mutation in genes encoding non-essential (glyco)proteins of the IBDV which still give rise to the production of antibodies in an infected host animal. A marker vaccine for Aujeszky's disease and companion diagnostic tests have proven their practical value in the control of this disease. Whereas such control programs for other viral infectious diseases in animals are under development, until the present invention a vaccine based on an IBDV vaccine strain which would fit in IBDV control programs has not been described yet.

[0009] Mundt and Köllner (Annual Meeting of the German Federal Research Institute for virus diseases in animals, March 1997)) disclose the construction of an IBDV mutant that fails to express a VP5 protein as a result of a genetically engineered mutation in the start codon of the VP5 gene. This mutation comprises the substitution of one nucleotide in the start codon of the VP5 gene. [0010] The present invention provides a birnavirus mutant as defined in the claims which is not able to produce a native VP5 protein as a result of a mutation in the VP5 gene of the birnavirus genome.

[0011] Preferably, the birnavirus mutant is an IBDV mutant or an IPNV mutant, the IBDV mutant being most preferred, in particular an IBDV mutant derived from a serotype 1 IBD virus is provided by the present invention.

[0012] It is demonstrated that an IBDV mutant that is not able to produce a VP5 protein is still able to infect poultry and to replicate in the infected host animals in vivo, i.e. evidence is provided that the gene encoding the VP5 protein is a non-essential gene. Example 3 and 4 show that VP5 IBDV can be re-isolated from organs of animals infected with the IBDV mutant and that the IBDV mutant induces a protective immune response in the infected animals.

[0013] Moreover, it has been established herein that part of the normal anti-IBDV immune response in poultry is directed to the VP5 region. This is rather surprising as the VP5 protein is considered to represent a non-structural viral protein (Mundt et al., J. Gen. Virol. 76, 437-443, 1995) and the immune response in an animal against a viral pathogen is usually elicited against the structural (glyco)proteins of the virus. These findings make the IBDV mutant and other birnavirus mutants according to the present invention a suitable vaccine candidate for a marker vaccine. Such a marker vaccine provides the possibility to determine whether animals are infected with a wild-type birnavirus, e.g. IBDV, or with a vaccine virus.

10

20

30

35

40

55

[0014] Additionally, it has been found that the VP5 protein is involved in the expression of virulence of the birnaviruses, in particular of IBDV, and that the inability of the virus mutants to produce the native VP5 protein leads to an attenuation

[0015] With the term "which is not able to produce a native VP5 protein" is meant that the birnavirus mutant produces a polypeptide that can be distinguished by serological tests from the native VP5 protein, or does not produce a VP5 protein at all. For example, in the former case, the birnavirus mutant produces only a fragment of the native birnavirus VP5 protein which lacks one or more immunogenic epitopes.

[0016] Preferably, the birnavirus mutant according to the invention produces no VP5 protein upon infection of a host cell.

25 [0017] As described above, the genomic organisation of the birnaviruses is well established: the IBDV and IPNV genome comprises a large segment A and a smaller segment B. The segment A of IBDV comprises a large open reading frame (ORF) encoding a polyprotein of about 110 kDa (VP2-VP4-VP3). The gene encoding the VP5 protein is identified in the prior art, and defined herein, as the small ORF on segment A of the birnavirus genome which precedes and partially overlaps the polyprotein encoding ORF (Bayliss et al., J. Gen. Virol. 71, 1303-1312, 1990; Spies et al., J. Gen. Virol. 71, 977-981, 1990; Havarstein L.S. et al., J. Gen. Virology 71, 299-308; 1990; Dobos et al., 1995, supra; Figures 1-3 herein and SEQ ID No.'s 1-7). The mutation introduced in the VP5 gene is such that it does not prevent the expression of the polyprotein.

[0018] SEQ ID No. 1 comprises the full length cDNA nucleotide sequence of segment B of IBDV strain P2, as well as the amino acid sequence of the VP1 protein encoded by segment B (see also SEQ ID. No. 2). SEQ ID No. 3 and 5 depict the full length cDNA sequence of segment A of IBDV strain D78 and the coding region of the VP5 protein and the polyprotein, respectively. SEQ ID 3 and 4 also show the amino acid sequence of the D78 VP5 protein. SEQ ID No. 5 and 6 show the amino acid sequence of the polyprotein VP2-VP4-VP3 of D78. SEQ ID No. 7 shows the 5'-end of segment A of strain D78, including the mutations introduced in the VP5 coding region. SEQ ID No. 8 shows the nucleotide sequence of segment B of strain D78 and the amino acid sequence of the D78 VP1 protein. The genomic organisation of both segments is also shown in Figure 1.

[0019] The ORF coding for VP5 is conserved in all hitherto published segment A sequences. The IBDV ORF encodes 145 amino acids resulting in a calculated molecular mass of 16.5 kDa. The nucleotide sequence of the ORF encoding the VP5 protein of IBDV strain D78 used herein is shown in SEQ ID No. 3 and 4. Natural variations may exist between individual IBDV isolates. These natural variations result from small differences in the genomes of these viruses. The nucleotide sequence of the segment A, including the nucleotide sequence of the VP5 gene for many IBDV isolates have been described in the prior art (Vakharia et al., Avian Diseases 36, 736-742, 1992; Bayliss et al., J. Gen. Virol. 71, 1303-1314, 1990; Hudson et al., Nuc. Acid Res. 14, 5001-5012, 1986; Schnitzler et al., J. Gen. Virol. 47, 1563-1571, 1993; Kibenge et al., J. Gen. Virol. 71, 569-577, 1990 and Virology 184, 437-440, 1991; Mundt et al., Virology 209, 10-18, 1995; Lana et al., Virus Genes 6, 247-259, 1992; Vakharia et al., Virus Res. 31, 265-273, 1994; Brown et al., Virus Res. 40, 1-15, 1996). The amino acid sequence of the VP5 protein from serotype I IBDV strains display a homology of at least 95% with the VP5 amino acid sequence shown in SEQ ID No. 3 and 4, whereas the homology between serotype II VP5 sequence and the amino acid sequence shown in SEQ ID No. 3 and 4 is at least 75%. Therefore, a preferred IBDV mutant according to the present invention is an IBDV mutant wherein the mutation is introduced in the VP5 gene having a homology of at least 75%, in particular at least 95% on the amino acid sequence level with the VP5 amino acid sequence shown herein.

[0020] Preferably an IBDV mutant according to the present invention is derived from any of the classical or variant (e.g. variant E or GLS) IBDV vaccine strains, such as those currently used in the field. Such suitable IBDV strains include the IBDV vaccine strains present in the commercially available vaccines: D78, PBG 98, LZ 228E, 89-03 (Intervet International B.V.), Bursine 2 (Fort Dodge Animal Health) and S 706 (Rhone Mérieux).

10

20

25

30

35

40

- [0021] A particular preferred IBDV mutant according to the invention is derived from the D78 strain comprising a VP5 gene encoding a protein having the amino acid sequence shown in SEQ ID No. 3 and 4.
- [0022] Alternatively, the parent birnavirus strain for the virus mutant according to the invention is a virulent birnavirus field strain. It is found herein that the VP5 protein is a factor associated with virulence, and that the absence of the native VP5 protein in a birnavirus results in an attenuated form of the virus.
- [0023] Preferably the invention provides a birnavirus mutant which is not able to produce a native VP5 protein as a result of a mutation in the part of the VP5 gene which does not overlap with the large ORF encoding the polyprotein.
- [0024] In particular, the birnavirus mutant according to the invention comprises a mutation in the 5'-end of the VP5 gene spanning nucleotides 1-30, preferably 1-20, more preferably 1-10. Most preferred is an birnavirus mutant having a mutation in nucleotides 1-3 of the VP5 gene.
- [0025] A mutation is understood to be a change of the genetic information in the VP5 gene with respect to the genetic information present in this region of the genome of naturally occurring birnavirus producing native VP5 protein. The mutation is, for example, a nucleic acid substitution, deletion, insertion or inversion, or a combination thereof.
- [0026] In a preferred embodiment of the present invention a birnavirus mutant is provided wherein the mutation is a substitution of one or more nucleotides. In particular, a nucleic acid substitution is introduced in the start codon, as a result of which the new codon encodes an amino acid different from methionine or represents a stop codon, preferably the nucleic acid substitution comprises at least two of the nucleotides of the start codon.
  - [0027] A further birnavirus mutant according to the invention comprises a substitution of one or more nucleotides in a codon(s) different from the start codon resulting in one or more stop codons, preferably in the 5'-end of the VP5 gene as defined above, if desired in addition to a substitution in the start codon as described above. Preferably, the birnavirus mutant comprises a stop codon in this region of the VP5 gene in each of the three reading frames.
  - [0028] Such a preferred birnavirus mutant may be an IBDV mutant having a mutation in the start codon, the fourth and the sixth codon of the VP5 gene, preferably resulting in the mutated codons shown in SEQ ID No. 7 and Figure 3.
  - [0029] Alternatively, a birnavirus mutant is provided wherein the mutation is a deletion. In particular, the deletion comprises less than 20, less than 10 or less than 5 nucleotides. Preferably, the deletion comprises a total number of nucleotides not dividable by three, resulting in a shift of the reading frame.
  - [0030] Preferably the deletion comprises one or more nucleotides of the start codon of the VP5 gene.
  - [0031] In an alternative embodiment of the present invention a bimavirus mutant is provided wherein the mutation comprises the insertion of a heterologous nucleic acid sequence in the birnavirus genome. A heterologous nucleic acid sequence is a nucleic acid sequence normally not present at the specific insertion site of the particular virus species. [0032] The heterologous nucleic sequence to be incorporated into the birnavirus genome is a nucleic acid fragment which either encodes a polypeptide or is a non-coding sequence. The nucleic acid fragment can be derived from any source, e.g. viral, eukaryotic, prokaryotic or synthetic, including oligonucleotides suitable for the interruption of the expression of the VP5 gene.
  - [0033] A suitable oligonucleotide for the interruption of the VP5 expression may comprise three translational stop codons in each of the possible reading frames in both directions, in addition to one or more appropriate restriction enzyme cleavage sites useful for the insertion of a second heterologous nucleic acid sequence. The length and nucleotide sequence of such a non-coding heterologous nucleic acid sequence is not critical, but preferably varies between 8-50 nucleotides.
  - [0034] In a further embodiment of the present invention a bimavirus mutant is provided which can be used not only for the preparation of a vaccine against infection by a specific birnavirus, but also against other poultry or fish infectious diseases. For example, a vector vaccine based on such an IBDV mutant offers the possibility to immunise against other avian pathogens by the expression of antigens of these avian pathogens within infected cells of the immunised host. Such an IBDV vector according to the present invention can be obtained by inserting a heterologous nucleic acid sequence encoding a polypeptide heterologous to the IBDV in the VP5 gene as defined herein.
  - [0035] The heterologous nucleic acid sequence may encode an antigen of an avian pathogen such as Newcastle disease virus, Infectious bronchitis virus, Marek's disease virus, avian encephalomyelitis virus, avian reovirus, avian influenza virus, chicken anaemia virus, Salmonella spp., E.coli, and Eimeria spp.
- [0036] Furthermore, an IBDV mutant according to the invention comprises in addition to the mutation in the VP5 gene, a mutation in the VP2 gene, wherein this gene expresses a chimeric protein comprising neutralising epitopes of more than one antigenic type of IBDV (e.g. classic, Variant-E and/or GLS). Preferably, such a mutant comprises the relevant protective VP2 epitopes of a variant GLS strain and classic strain. In particular, the mutated VP2 gene is a GLS VP2 gene comprising a nucleic acid sequence fragment encoding the B69 epitope. The construction of such a mutated VP2 genes is described in Snyder et al., Avian Diseases 38, 701-707, 1994. Furthermore, nucleic acid sequences encoding polypeptides for pharmaceutical or diagnostic applications, in particular immuno-modulators such as lymphokines, interferons or cytokines, may be incorporated into the VP5 gene. The heterologous nucleic acid sequence may also encode a screenable marker, such as E. coli β-galactosidase or E. coli β-glucuronidase.

[0037] The construction of birnavirus mutants, in particular of IBDV mutants according to the present invention can be achieved by means of the recently established infectious cRNA system for IBDV (Mundt and Vakharia, Proc. Natl. Acad. Sci. USA 93, 11131-11136, 1996). This reverse genetics system opens the possibility to introduce mutations in the RNA genome of the IBD virus, in particular in the VP5 gene. The most important step in this reverse genetics system is to provide full length cDNA clones of the segments A and B of IBD virus. cDNA constructs comprising the segment A or B, including the nucleotides of the 5'- and 3'- ends of both these segments can be generated according to the method described by Mundt and Vakharia (1996, supra). Additionally, these constructs comprise a RNA polymerase promoter operably linked to either of the segments. The promoter can be the promoter for the T7, SP6 or T3 polymerase, the T7 promoter being preferred. Mutations can be introduced into the VP5 gene by means of methods generally known in the art for this purpose. In particular, the mutation(s) are introduced by means of site directed mutagenesis.

5

10

15

20

30

35

45

50

55

[0038] For example, in a first step a cDNA fragment is provided comprising at least a substantial part of the VP5 gene. In the next step suitable primer pairs are designed and hybridised with the VP5 sequence containing fragment. The 5'-primer comprises in addition to sequences complementary to the VP5 sequence, nucleotides which harbour the desired mutation, e.g. a mutation which changes the ATG start codon to an AGG (arginine) codon. Moreover, the 5'-primer is provided with an upstream nucleotide sequence representing a suitable restriction enzyme cleavage site which allows the restoring of the complete 5'-end non-coding sequence. Subsequently, the new mutated fragment is amplified using PCR and the new fragment is introduced in the starting sequence by replacing the native nucleic acid sequence using appropriate restriction enzymes. In the next step plus-sense transcripts of the segment A and B are generated in vitro with (T7) RNA polymerease, after which the synthetic transcripts are purified using conventional RNA purification techniques. The recombinant IBDV mutant according to the invention is obtained after transfection of suitable cells (e.g. VERO cells, QM-7 cells or CEC cells) with the synthetic RNA transcripts of both segments of the IBDV genome, if desired in the presence of transfection-enhancing compositions, such as Lipofectin. Finally the recombinant IBDV is harvested from the supernatant of the transformed cells.

[0039] Methods for introducing a mutation in the birnavirus genome are described herein, but are also generally used in the art (Mundt and Vakharia, 1996, supra; Current Protocols in Molecular Biology, eds.: F. M. Ausubel et al., Wiley N.Y., 1995 edition, pages 8.5.1.-8.5.9.)

[0040] Further to the unexpected finding by the present inventors that the VP5 ORF of IBDV is a non-essential region of the IBDV genome, it has also been found that an IBDV mutant according to the present invention is able to induce a protective immune response, i.e. animals immunised with a vaccine comprising the IBDV mutant are protected against virulent challenge. Moreover, it has been found that anti-sera of animals infected with naturally occurring IBDV comprise antibodies directed to the non-structural VP5 protein and that these antisera can be distinguished from anti-sera derived from animals infected with an IBDV mutant according to the present invention. In addition, it has been found that the IBDV mutant as described above is attenuated if compared with the parent IBD virus which is able to produce the native VP5 protein.

[0041] Therefore, another aspect of this invention is a vaccine for use in the protection of animals against birnavirus infection comprising the birnavirus mutant as characterised above, together with a pharmaceutical acceptable carrier or diluent. In particular, the vaccine according to the invention is a vaccine for use in the protection of poultry against infectious bursal disease comprising the IBDV mutant described above.

[0042] The birnavirus mutant according to the present invention can be incorporated into the vaccine as live or inactivated virus.

**[0043]** A vaccine according to the invention can be prepared by conventional methods such as for example commonly used for the commercially available live- and inactivated IBDV vaccines. Briefly, a susceptible substrate is inoculated with an IBDV mutant according to the invention and propagated until the virus replicated to a desired infectious titre after which IBDV containing material is harvested.

[0044] Every substrate which is able to support the replication of IBD viruses can be used in the present invention, including primary (avian) cell cultures, such as chicken embryo fibroblast cells (CEF) or chicken kidney cells (CK), mammalian cell lines such as the VERO cell line or the BGM-70 cell line, or avian cell lines such as QT-35, QM-7 or LMH. Usually, after inoculation of the cells, the virus is propagated for 3-10 days, after which the cell culture supernatant is harvested, and if desired filtered or centrifuged in order to remove cell debris.

[0045] Alternatively, the IBDV mutant is propagated in embryonated chicken eggs. In particular, the substrate on which these IBD viruses are propagated are SPF embryonated eggs. Embryonated eggs can be inoculated with, for example 0.2 ml IBDV mutant containing suspension or homogenate comprising at least  $10^2$  TCID $_{50}$  per egg, and subsequently incubated at 37 °C. After about 2-5 days the IBD virus product can be harvested by collecting the embryo's and/or the membranes and/or the allantoic fluid followed by appropriate homogenising of this material. The homogenate can be centrifuged thereafter for 10 min at 2500 x g followed by filtering the supernatant through a filter (100  $\mu$ m).

[0046] The vaccine according to the invention containing the live virus can be prepared and marketed in the form of a suspension or in a lyophilised form and additionally contains a pharmaceutically acceptable carrier or diluent cus-

tomary used for such compositions. Carriers include stabilisers, preservatives and buffers. Suitable stabilisers are, for example SPGA, carbohydrates (such as sorbitol, mannitol, starch, sucrose, dextran, glutamate or glucose), proteins (such as dried milk serum, albumin or casein) or degradation products thereof. Suitable buffers are for example alkali metal phosphates. Suitable preservatives are thimerosal, merthiolate and gentamicin. Diluents include water, aqueous buffer (such as buffered saline), alcohols and polyols (such as glycerol).

[0047] If desired, the live vaccines according to the invention may contain an adjuvant. Examples of suitable compounds and compositions with adjuvant activity are the same as mentioned below.

[0048] Although administration by injection, e.g. intramuscular, subcutaneous of the live vaccine according to the present invention is possible, the vaccine is preferably administered by the inexpensive mass application techniques commonly used for IBDV vaccination. For IBDV vaccination these techniques include drinking water and spray vaccination.

10

15

20

25

30

40

45

50

[0049] Alternative methods for the administration of the live vaccine include in ovo, eye drop and beak dipping administration.

[0050] In another aspect of the present invention a vaccine is provided comprising the birnavirus mutant in an inactivated form. The major advantage of an inactivated vaccine is the extremely high levels of protective antibodies of long duration that can be achieved.

[0051] The aim of inactivation of the viruses harvested after the propagation step is to eliminate reproduction of the viruses. In general, this can be achieved by chemical or physical means. Chemical inactivation can be effected by treating the viruses with, for example, enzymes, formaldehyde,  $\beta$ -propiolactone, ethylene-imine or a derivative thereof. If necessary, the inactivating compound is neutralised afterwards. Material inactivated with formaldehyde can, for example, be neutralised with thiosulphate. Physical inactivation can preferably be carried out by subjecting the viruses to energy-rich radiation, such as UV light or  $\gamma$ -rays. If desired, after treatment the pH can be adjusted to a value of about 7.

[0052] A vaccine containing the inactivated birnavirus mutant can, for example comprise one or more of the abovementioned pharmaceutically acceptable carriers or diluents suited for this purpose.

[0053] Preferably, an inactivated vaccine according to the invention comprises one or more compounds with adjuvant activity. Suitable compounds or compositions for this purpose include aluminium hydroxide, -phosphate or -oxide, oil-in-water or water-in-oil emulsion based on, for example a mineral oil, such as Bayol F® or Marcol 52® or a vegetable oil such as vitamin E acetate, and saponins.

[0054] The vaccine according to the invention comprises an effective dosage of the birnavirus mutant as the active component, i.e. an amount of immunising birnavirus material that will induce immunity in the vaccinated birds against challenge by a virulent virus. Immunity is defined herein as the induction of a significant higher level of protection in a population of birds after vaccination compared to an unvaccinated group.

**[0055]** Typically, the live vaccine according to the invention can be administered in a dose of  $10^2$ - $10^9$  TCID<sub>50</sub> infectious dose<sub>50</sub> (TCID<sub>50</sub>) per animal, preferably in a dose ranging from  $10^{5.0}$ - $10^{7.0}$  TCID<sub>50</sub>, and an inactivated vaccines may contain the antigenic equivalent of  $10^5$ - $10^9$  TCID<sub>50</sub> per animal.

[0056] Inactivated vaccines are usually administered parenterally, e.g. intramuscularly or subcutaneously.

[0057] Although, the IBDV vaccine according to the present invention may be used effectively in chickens, also other poultry such as turkeys, guinea fowl and partridges may be successfully vaccinated with the vaccine. Chickens include broilers, reproduction stock and laying stock.

[0058] The age of the animals receiving a live or inactivated vaccine according to the invention is the same as that of the animals receiving the conventional live- or inactivated IBDV vaccines. For example, broilers (free of maternally derived antibodies-MDA) may be vaccinated at one-day-old, whereas broilers with high levels of MDA are preferably vaccinated at 2-3 weeks of age. Laying stock or reproduction stock with low levels of MDA may be vaccinated at 1-10 days of age followed by booster vaccinations with inactivated vaccine on 6-8 and 16-20 weeks of age.

[0059] The invention also includes combination vaccines comprising, in addition to the IBDV or IPNV mutant according to the invention, one or more immunogens derived from other pathogens infectious to poultry or fish, respectively. [0060] Preferably, the combination vaccine additionally comprises one or more vaccine strains of infectious bronchitis virus (IBV), Newcastle disease virus (NDV), egg drop syndrome (EDS) virus, turkey rhinotracheitis virus (TRTV) or reovirus.

[0061] In addition to a marker vaccine for birnaviruses, the availability of an appropriate diagnostic test is an essential requirement for the application of a birnavirus eradication control programme. Such a diagnostic test is provided herein and comprises a method for determining IBDV infection in poultry and IPNV infection in fish, i.e. it provides a method for distinguishing an animal in the field vaccinated with a vaccine as described above, from an animal infected with a naturally-occurring IBDV or IPNV.

[0062] Therefore, the present invention provides a method for the detection of birnavirus infection, in particular for the detection of IBDV infection in an animal comprising the step of examining a sample of the animal for the presence of VP5 antibodies or antigens. The animal is an animal from the field and is in particular an avian species, preferably

a chicken. The sample coming from the animal may be any sample in which IBDV antibodies or antigens are present, e.g. a blood, serum or tissue sample, the serum sample being preferred.

[0063] A preferred method for determining birnavirus infection in an animal is a method for the detection of antibodies against the VP5 protein, comprising the steps of:

- (i) incubating a sample suspected of containing anti-birnavirus antibodies, with VP5 antigen,
- (ii) allowing the formation of antibody-antigen complex, and
- (ii) detecting the presence of the antibody-antigen complex.

[0064] The design of this immunoassay may vary. For example, the immunoassay may be based upon competition or direct reaction. Furthermore, protocols may use solid supports or may use cellular material. The detection of the antibody-antigen complex may involve the use of labelled antibodies; the labels may be, for example, enzymes, fluorescent-, chemiluminescent-, radio-active- or dye molecules.

[0065] Suitable methods for the detection of the VP5 antibodies in the sample include the enzyme-linked immunosorbent assay (ELISA), immunofluorescent test (IFT) and Western blot analysis.

[0066] In an exemplifying ELISA, the wells of a polystyrene micro-titration plate are coated with VP5 antigen. Next, the wells of the coated plates are filled with chicken serum and serial dilutions are made. After incubation, chicken anti-VP5 protein serum antibodies are determined by detecting antibody (monoclonal or polyclonal) with the same specificity as the coated one, but which is labelled (e.g. with biotin). The labelled antibody will occupy the free antigens that have not been occupied by anti-VP5 antibodies in the chicken serum. For example, horse radish peroxidase coupled to avidin may be added and the amount of peroxidase is measured by an enzymatic reaction. If no antibodies against VP5 are present in the chicken serum sample then a maximum absorption is obtained. If the serum contains many antibodies against VP5 then a low absorption is expected. Alternatively, after the incubation with chicken serum, the amount of antibodies present in the serum that bound to the VP5 antigen may be determined directly by using an anti-chicken conjugate followed by the enzymatic reaction.

[0067] In a sandwich ELISA the wells of a polystyrene micro-titration plate can be coated with a monoclonal antibody directed against the VP5 protein. Next, the wells of these coated plates are incubated with VP5 antigen. After the antigen is captured, the wells are filled with the chicken serum and serial dilutions are made. Subsequently, the protocol as described above may be followed. This test can also be carried out by using polyclonal serum against VP5 instead of the coated monoclonal antibodies.

[0068] In another diagnostic test (Western blot analysis), the VP5 antigen (containing) material is subjected to SDS-PAGE. Next, the separated proteins are electroblotted onto nitro-cellulose membrane. Thereafter, the membranes can be cut into lanes and the lanes are incubated with the chicken serum. The presence of VP5 antibodies in the sample can be determined by examination whether antibodies bound to the VP5 antigen, for example by using an anti-chicken conjugate followed by an enzymatic reaction. If antibodies against VP5 are present then a band at about 17 kDa is identifiable.

[0069] The VP5 antigen may be any VP5 protein (fragment) comprising material which allows the formation of the VP5 antigen-VP5 antibody complex. Preferably, the VP5 antigen comprises the expression product of a conventional recombinant host cell or virus, e.g. such as E.coli expressed VP5 (Mundt et al., J. Gen. Virol. 76, 437-443, 1995) or baculovirus expressed protein (Vakharia et al., Vaccine 12, 452-456, 1994; Vakharia et al., J. Gen Virol. 74, 1201-1206, 1993). In a further embodiment of the present invention a diagnostic test kit is provided which is suitable for performing the diagnostic test according to the invention as described above.

[0070] In particular, a diagnostic test kit is provided which comprises in addition to the components usually present, the VP5 antigen (if desired coated onto a solid phase) as the immunological reagent. Other components usually present in such a test kit include, biotin or horseradish peroxidase conjugated antibodies, enzyme substrate, washing buffer etc.

[0071] To determine birnavirus VP5 antigen in a test sample from an animal in the field, VP5-specific antibodies are used as the immunological reagent, preferably fixed to a solid phase. The test sample is added, and after an incubation time allowing formation of the antibody-antigen complex, a second labelled antibody may be added to detect the complex.

50

5

10

15

20

25

30

35

40

45

#### **EXAMPLES**

#### Example 1.

10

15

20

25

30

35

40

45

50

55

5 Construction and analysis of recombinant VP5 IBD virus

#### Construction of full length VP5 clone of IBDV segment A.

[0072] To construct a VP5-negative IBDV, the *EcoRI* site immediately following the 3'-end of the full length cDNA of strain D78 segment A (pUC19FLAD78; Mundt and Vakharia, Proc. Natl. Acad. Sci. USA <u>93</u>, 11131-11136, 1996) was deleted. An *EcoRI* - *KpnI* fragment containing the T7 polymerase binding site followed by the complete segment A sequence was excised and inserted into *EcoRI* - *KpnI* cleaved vector pUC18 after inactivation of the unique *NdeI* within the vector sequence resulting in plasmid pAD78/EK. Thereafter, the genomic region encompassing the initiation codon for VP5 was amplified in two pieces using primers A1F5' and VP5MutR, and VP5MutF and A2R, respectively (see Table 1 for sequence and location of primers). PCR fragments were cloned separately and were subsequently fused via a unique *AfIII* site which had been created by mutations within respective primers (see Fig. 2). An *EcoRI* - *NdeI* fragment containing the T7 polymerase binding site, and the 5'-part of segment A including the introduced mutations was excised and used to substitute the wild-type *EcoRI* - *NdeI* fragment in pAD78/EK to yield plasmid pAD78/VP5-. Of the three mutations introduced one altered the initiation methionine codon for VP5 into an arginine codon (Fig. 2).

<u>Table 1</u>: Sequence of oligonucleotide primers used for generating mutant constructs.

<sup>a</sup> Nucleotide sequence	Orientation	Designation .	Nucleotide no.
AGAGAATTC <i>TAATACGACTCACTATA</i> GGA	+	A1F5'	1-18
TACGATCGGTCTGAC			
TGGGCCTGTCACTGTCACATGT	-	A2R	716 - 740
CATTGCTCTGCAGTGTGTGAGC	-	A3R	338 - 362
CTACAACGCTATCCTTAACGGTTAGTA	+	VP5MutF	80 - 109
GAG			
CTCTACTAACCCTTAAGGATAGCGTTGT		VP5MutR	80 - 109
AG			

a) Underlined nucleotides denote virus specific nucleotides. T7 promotor sequences are marked in italics. Mutated nucleotides are bold and orientation of the primer is shown for sense (+) and antisense (-). Primer positions are given according to the published sequence of serotype I strain P2 (Mundt et al., Virology 209, 209-218, 1995).

[0073] Virus recovery from cRNA. For *in vitro* transcription of RNA plasmids pAD78/EK, pAD78/VP5- and pBP2 (Fig. 2) were linearized by cleavage with *Bsr*Gl and *Pst*I, respectively. Treatment of linearized DNA, transcription and purification of RNA, and transfection were carried out as described by Mundt and Vakharia (1996, supra) with the exception that secondary CEC were used for the transfection experiments. Three days after transfection a CPE was visible in CEC. Cells were freeze/thawed, centrifuged at 700 x g to eliminate cellular debris, and the resulting supernatants were filtrated through 0.45 µm filters and stored at -20°C. For the transfection experiments full length cDNA clones of segment A of strain D78 capable of expressing (pAD78/EK) or unable to express VP5 (pAD78/VP5-) were transcribed into synthetic RNA and cotransfected with segment B full length cRNA into CEC. Resulting virus progeny IBDV/EK and IBDV/VP5- was further characterised.

[0074] Analysis of transfection progeny by immunofluorescence and Radioimmunoprecipitation assay

(RIPA). VP5 was expressed in E.coli as described in Mundt et al. (J. Gen. Virol. <u>76</u>, 437-443, 1995). Rabbit monospecific polyclonal anti serum and mouse monoclonal antibodies against VP5 were prepared according to standard protocols. Vero cells infected with IBDV/VP5-, IBDV/EK, and non-infected cells , respectively, were incubated with rabbit anti-IBDV serum, rabbit anti-VP5 serum and with anti-VP5 mAb DIE 7, and stained with fluoresceine-conjugated secondary antibodies. Both antisera and the monoclonal antibody recognised IBDV antigens in the cytoplasm of IBDV/EK infected cells. In contrast, whereas the anti-IBDV serum readily detected viral antigens in IBDV/VP5- infected cells, neither the monospecific anti VP5-serum nor the monoclonal anti-VP5 antibody exhibited specific reactivity. None of these immunological reagents reacted with non-infected controls.

[0075] To analyse viral proteins expressed during replication lysates of radioactively labelled CEC infected with IBDV/VP5- (Fig 4, lanes 1-3) and IBDV/EK (Fig. 4, lanes 4-6) were immunoprecipitated with rabbit anti-IBDV serum, rabbit anti-VP5 serum and mAb DIE 7. Non-infected CEC were used as control (Fig. 4, lanes 7-9). IBDV/EK (lane 4) as well as IBDV/VP5- (lane 1) infected CEC showed viral proteins VP2, VP3, and VP4 after precipitation with rabbit anti-IBDV serum. The rabbit anti-VP5 serum (lane 5) and mAb DIE 7 (lane 6) precipitated VP5 with a molecular mass of 21 kDa only from IBDV/EK infected cells. No specific reactivity was detectable in IBDV/VP5- infected CEC after precipitation with rabbit-anti-VP5 (lane 2) as well as the VP5 specific mAb DIE 7 (lane 3). Non-infected CEC showed no specific reactivity (lanes 7-9).

[0076] Replication of IBDV/VP5<sup>-</sup> in CEC. To assay replication of IBDV/VP5<sup>-</sup> in more detail one step growth was analysed (Fig. 5). Confluent secondary CEC were infected with IBDV/EK and IBDV/VP5<sup>-</sup> with 10<sup>72</sup> TCID<sub>50</sub>, respectively. Immediately after overlaying the infected cells with 5 ml growth medium, supernatant from one infected CEC tissue plate of each virus was removed and stored at -20°C (0 h p.i.). Remaining tissue culture plates were further incubated and 4h, 8h, 16h, 24h, and 48h p.i. supernatants were removed and stored at -20°C. Supernatants were centrifuged and titrated according to standard methods. The TCID<sub>50</sub> at the different time points after infection showed that the VP5 expressing virus (IBDV/EK) replicated faster than the virus mutant lacking VP5 (IBDV/VP5<sup>-</sup>). 16 h after infection IBDV/EK showed a 100-fold higher than IBDV/VP5<sup>-</sup> (Fig. 5). However, at 48 h p.i. IBDV/VP5<sup>-</sup> reached a titre of 10<sup>7.2</sup> TCID<sub>50</sub>/ ml which was similar to IBDV/EK (10 <sup>7.45</sup>/ml)

[0077] Preparation of recombinant IBDV VP5-2. Plasmid pAD78/VP5-2 was prepared by techniques similar to those described above. The nucleotide sequence of part of the mutated VP5 gene is shown in SEQ ID No. 7 and Figure 3. A restriction enzyme fragment harbouring the mutations was used to substitute the wild-type EcoRI - Ndel fragment in pAD78/EK. An outline of the protocol for the preparation of the recombinant plasmid is shown in Figure 3. The organisation of pBD78 is also depicted in Figure 3. The recombinant virus was prepared as described above, except for the fact that segment B of strain D78 (SEQ ID No. 8) was used and QM-7 cells were used for the transfection experiment.

### Example 2

### Identification of VP5 protein in different IBDV strains

[0078] Different strains of IBDV were investigated for the expression of the VP5-gene. This was done by making use of the immuno-fluorescence technique (IFT). Chicken embryo fibroblasts grown in microtiterplates were infected with different IBDV strains. Three to 5 days after incubation at 37°C cells were fixed with 70% ethanol, then treated with polyclonal rabbit anti IBDV serum (R1928), polyclonal rabbit anti VP5 serum (RαVP5) or monoclonal antibody directed against VP5 (DIE7), respectively. Binding of the poly- or monoclonal antibodies to the different IBDV strains was visualised by making use of a fluorescence labelled conjugate (goat-anti-rabbit or goat-anti-mouse). The results are shown in Table 2:

Table 2

Identification of differen	ent sero- and subtypes	of IBDV strains. Determ	nination of the p	resence of VP	5 proteins.
IBDV-serotype	IBDV-subtype	IBDV-strain	R1928	RαVP5	DIE7
I	Classical	D78	+	+	+
ı	Classical	228TC	+	+	+
	Classical	PBG98	+	+	+
ı	Classical	Ram0404	+	+	+
, '	Classical	IBDV/EK	+	+	+
	Classical	IBDV/VP5-	+	•	-

45

5

10

15

20

25

35

40

50

5**5** 

Table 2 (continued)

Identification of differen	ent sero- and subtypes	of IBDV strains. Deter	mination of the p	resence of VF	5 proteins.
IBDV-serotype	IBDV-subtype	IBDV-strain	R1928	RαVP5	DIE7
1	GLS	GLS	+	+	+
ı	Variant-E	8903	+	+	+
II	TY89	TY89	+	+	+

**[0079]** From these data it can be concluded that the different strains of IBDV belonging to different sero- and subtypes do express the VP5-gene. Furthermore, the recombinant VP5- IBDV vaccine strain can be differentiated from field and vaccine viruses, thereby enabling the recombinant VP5- virus to be used as a marker vaccine.

### Example 3

In vivo testing of the recombinant VP5+ and VP5- IBDV vaccines in comparison with a commercial available live IBDV vaccine.

[0080] Preparation of IBDV vaccine. Primary chicken embryo fibroblast (CEF) cells were prepared at a final concentration of 2x10<sup>6</sup>/ml. The cells were cultured in Eagles minimum essential medium containing 5% fetal calf serum. To 25 ml of this cell suspension 0.1 ml IBDV/EK or IBDV/VP5 virus (having an infectious titre of about 3.0 log10 TCID<sub>50</sub>/ml) was added. After incubation for 5 days in a high-humidity incubator at 37°C, the total suspension was used in the animal experiment without further purification. The infectious titre of the supernantant was 10<sup>7.1</sup> TCID50/ml.

[0081] Animal experiment. In this study the potency of different vaccines (VP5 positive strain IBDV/EK and a VP5 negative strain IBDV/VP5<sup>-</sup>, and the commercial available IBDV vaccine Nobilis strain D78, Intervet International B.V., NL) was investigated. SPF chicks of 3 weeks old were treated as indicated in the treatment schedule. Treatment Schedule:

Days after vaccination		Groups		
	1	2	3	4
00	IBDV/EK	IBDV/VP5	D78	-
03	x	x1	х	×
07	x,bl	x1,bl	x,b	x,bl
14	x,bl	x,bl	x,bl	x,bl
20	x,bl	x,bl	x,bl	x,bl
21	ch	ch	ch	ch
24	×	×	х	х
31	+	+	+	+

VP5<sup>+</sup> Bursal disease vaccination with VP5 positive vaccine clone, eye-drop route, dose 10<sup>46</sup> TCID<sub>50</sub>/animal, 0.1 ml/animal.

VP5<sup>-</sup> Bursal disease vaccination with VP5 negative vaccine clone, eye-drop route, dose 10<sup>59</sup> TCID<sub>50</sub>/animal, 0.1 ml/animal.

D78 Bursal disease vaccination with IBDV VACCINE NOBILIS STRAIN D78, eye-drop route, one field dose.

ch Challenge with Bursal disease virus, Farragher strain F52/70, eye-drop route, dose 10<sup>2.0</sup> CID<sub>50</sub>/animal, 0.1 ml/animal.

bl Serological examination; VN-test and/or Western blotting.

- x Histological examination (H.E. staining) and MCA-8 ELISA on bursae.
- x1 Histological examination (H.E. staining) and MCA-8 ELISA on bursae and reisolation of virus from bursa of Fabricius.
- + Clinical examination and after 10 days histological examination of the bursa.

15

20

25

10

5

35

30

45

40

55

50

#### Detection of virus in the bursa of Fabricius.

5

10

15

20

25

30

35

40

[0082] Three, 7, 14 and 20 days after eye-drop vaccination, animals were sacrificed and blood and bursae obtained. The presence of virus in the bursa was determined with an enzyme-linked immunosorbent assay (ELISA) making use of the monoclonal antibody 8 (MAB-8). MAB-8 is directed specifically against IBDV. Data are depicted in Table 3.

[0083] Furthermore, 3 and 7 days after vaccination, bursae from animals of group 2 were investigated for the presence of the recombinant VP5- virus. For that purpose bursae were homogenised and cultured on chicken embryo fibroblasts. The presence of the VP5- virus was determined by IFT using polyclonal rabbit sera against IBDV or VP5 or monoclonal antibodies against VP5. From 13 out of 15 bursae (87%) investigated, VP5- virus could be reisolated and identified (positive for R1928 and negative for RαVP5 and DIE7). This indicates that the virus upon animal passage is still VP5-, indicating that the virus is stable and does not revert to VP5+. Furthermore, by using the different polyand monoclonal antibodies VP5- vaccine virus can be discriminated from all other vaccine and/or field IBDV viruses. Therefore, the VP5- vaccine may be used as a marker vaccine.

[0084] Three days after challenge no virus could be detected in groups 1, 2 and 3 with the MCA-8 ELISA. In contrast, all animals of group 4 (non-vaccinated control group) contained challenge virus in the bursa of Fabricius, 3 days after challenge. The results show that animals vaccinated with recombinant VP5+ (group 1), recombinant VP5- (group 2) and IBDV vaccine Nobilis D78 (group 3) were protected against severe challenge.

Table 3

			or chall		A-8 ELISA at different	radys and vaccinati
	Da	ys after vacc	ination→		Days after challenge	
	3	7	14	20	3	1
Group↓		Virus detection by ELISA↓				Protection↓
1 VP5+	2/8*	1/7	0/2	0/3	0/5	100%
2 VP5-	0/8	0/7	0/2	0/3	0/5	100%
3 D78	1/8	6/7	0/2	0/3	0/5	100%
4 -	0/8	0/7	0/2	0/3	5/5	0%

\*Number of positive bursae per total number tested.

#### Detection of lesions in the bursa of Fabricius.

[0085] The microscopic average lesion score induced by the different IBDV (recombinant) vaccines or the challenge virus are depicted in Table 4.

[0086] Before challenge, animals vaccinated with the recombinant VP5<sup>+</sup> IBDV vaccine (group 1) or vaccinated with IBDV vaccine Nobilis D78 (group 3) showed mild to moderate lesions in the bursa. Three days after challenge only chronic lesions were observed in the bursa of Fabricius, indicating that the animals of groups 1 and 3 were protected against challenge. Furthermore, 10 days after challenge only very mild lesions (0-20% lymphocytic depletion) were observed in the bursa of the animals vaccinated with VP5<sup>+</sup> recombinant IBDV vaccine or with Nobilis vaccine D78. In contrast animals not vaccinated and challenged showed severe lesions 10 days after challenge. In other words all animals (100%) of groups 1 and 3, vaccinated with the VP5<sup>+</sup> recombinant IBDV vaccine or with Nobilis vaccine D78 were protected against severe challenge.

[0087] Three, 7, 14 and 20 days after vaccination and 3 and 10 days after challenge with the recombinant VP5- IBDV vaccine, animals of group 2 showed no to hardly any lesions (0-20% lymphocytic depletion) in the bursa. All animals of group 2, vaccinated with the VP5- recombinant IBDV vaccine, were protected against severe challenge. When animals vaccinated with the recombinant VP5- IBDV vaccine are compared to animals of groups 1 or 3 (vaccinated with a recombinant VP5+ or commercial available vaccine) the recombinant VP5- vaccine induces less lesions and therefore, is safer, milder than the vaccines tested in this experiment.

[0088] Three days post-challenge, all non-vaccinated animals of group 4 showed severe acute lesions in the bursa (total lymphocyte depletion, score 5.0). Ten days after challenge, all animals (17 out of 17 animals) showed total lymphocytic depletion, indicating that these animals were not protected against severe challenge. Animals that died after challenge, all showed severe lesions in the bursa of Fabricius. It was concluded that control group 4 was not protected against severe challenge indicating that the test conditions were optimal.

#### Table 4

Average bursal lesion score at different days after vaccination or challenge. The average lesion score is calculated as follows: all lesion scores from the animals per group on a certain day are added. This number is then divided by the total number of animals investigated in that group on that day. Individual scores range from 1 to 5. Score 0 = no lymphocytic depletion, score 1 = 0 - 20%; score 2 = 20 - 40%; score 3 = 40 - 60%; score 4 = 60 - 80% and score 5 = 80 - 100 % lymphocytic depletion (total lymphocytic depletion).

		Days after v	vaccination→		Days after	challenge→	
	3	7	14	20	3	10	
Group↓			Bursal le	sions score↓		<u> </u>	Protection↓
1 VP5+	0.8	2.9	1.0	1.0	1.0°	0.6	100%
2 VP5-	0.0	0.0	0.5	0.0	0.0c	0.1	100%
3 D78	0.1	2.4	3.5	2.0	2.8c	1.1	100%
4 -	0.0	0.0	0.0	0.0	5.0a	5.0	0%

a Acute lesions

5

10

15

20

25

30

35

40

45

50

#### Serological response.

[0089] The serological response of the animals was determined by measuring the ability of blood serum to neutralise a classical infectious bursal disease virus strain in a virus neutralising (VN) test. Serum was investigated 3, 7, 14 and 20 days after vaccination. The average neutralising titres are shown in Table 5.

[0090] The results show that recombinant IBDV vaccine VP5+ applied to chickens of group 1 induced a good and high serological response 20 days after vaccination which is comparable to the serological response of the chickens vaccinated with the commercial IBDV vaccine Nobilis strain D78 (group 3). The recombinant IBDV vaccine VP5- applied to chickens of group 2 induced also a good serological response. A titre of 9.4 log2 was observed 20 days after vaccination. The serological response induced by the recombinant VP5- IBDV vaccine was delayed when compared to the serological response induced by the recombinant IBDV VP5+ vaccine or the commercial IBDV vaccine Nobilis strain D78.

[0091] The non-vaccinated group 4 showed no serological response to IBDV.

Table 5

Average IBDV-VI	N-titres for groups 1 to 4	at different days after	vaccination, expressed	as log2 of the dilution.
Group		Days after	vaccination	
	3	. 7	14	20
1 VP5+	≤ 1.0 ± 0.0	7.1 ± 1.7	10.2 ± 1.4	11.9 ± 1.8
2 VP5-	≤ 1.0 ± 0.0	2.1 ± 1.7	6.3 ± 2.9	9.4 ± 1.4
3 D78	´ ≤ 1.0 ± 0.0	5.2 ± 2.8	10.3 ± 1.3	11.6 ± 1.5
4-	≤ 1.0 ± 0.0	≤ 1.0 ± 0.0	≤ 1.0 ± 0.0	≤ 1.0 ± 0.0

### Serological differentiation between antisera.

[0092] The serological response against VP5 was investigated by making use of western blot analysis. For this purpose the VP5 protein was expressed in the E. coli or baculo expression system. The expressed proteins were separated by SDS PAGE. Next the proteins were electroblotted onto a nitro-cellulose membrane. Thereafter, the membrane was cut into lanes and the lanes were incubated with rabbit anti-VP5 serum, chicken serum directed against VP5+ recombinant vaccine, chicken serum directed against VP5+ recombinant vaccine or negative serum from SPF chickens. Data are summarised in Table 6. As can be seen from Table 6, the VP5- serum does not induce a serological response against VP5. In contrast the rabbit anti-VP5 serum and chicken serum directed against VP5+ recombinant vaccine do recognise the VP5-protein and thus induces a serological response against VP5. This indicates that chicken serum may be used to investigate if animals are exposed to a virus that expresses the VP5 protein (e.g. field virus) or

<sup>&</sup>lt;sup>C</sup> Chronic lesions

to the VP5 recombinant vaccine.

Table 6

Western blot analysis. Serum from animals vaccinated with VP5+ or VP5-re chicken serum and anti VP5-rabbit serum were investigated for their research.	
Identification of serum sample	Immuno-blot
VP5+ vaccinated animal, serum sample 20d after vaccination	positive
VP5- vaccinated animal, serum sample 20d after vaccination	negative
Non-vaccinated control, serum sample at 20d	negative
Rabbit anti VP5 serum	positive

#### Mortality and clinical signs.

[0093] None of the animals vaccinated with VP5+ IBDV vaccine (group 1), vaccinated with recombinant VP5- IBDV vaccine (group 2) or vaccinated with the commercial IBDV vaccine Nobilis strain D78 (group 3), died or showed clinical signs of infectious bursal disease after challenge, indicating that the animals were protected against severe challenge. All animals in the non-vaccinated control group were not protected against severe challenge.

### Example 4

5

10

15

20

25

30

35

40

45

#### In vivo testing of the recombinant VP5-2 vaccine

[0094] Preparation of the IBDV vaccines. Primary chicken embryo fibroblasts (CEF) cells were prepared at a final concentration of 2 x  $10^6$ /ml. The cells were cultured in Eagles minimum essential medium containing 5% fetal calf serum. To 15 ml of this cell suspension 0.1 ml IBDV/VP5-2 (D78/D78/VP5-) virus was added. After incubation for 6 days in a high humidity incubator at 37°C, the supernatant was titrated. The infectious titre of the supernatant was  $10^{8.2}$  TCID<sub>50</sub>/ml. For the second animal experiment the supernatant was diluted to result in a vaccine dose of  $10^{5.5}$  TCID<sub>50</sub>/animal and for the first animal experiment the supernatant was diluted to result in a vaccine dose of  $10^{4.0}$  TCID<sub>50</sub>/animal or  $10^{5.0}$  TCID<sub>50</sub>/egg.

[0095] First animal experiment. The effect of the vaccine is assessed by measurement of the serological response and resistance to challenge obtained from administering a challenge virus at the age of 14 days. The vaccine (10<sup>5.0</sup> TCID<sub>50</sub>/egg or 10<sup>4.0</sup> TCID<sub>50</sub>/animal of D78/D78/VP5<sup>-</sup>) was applied *in ovo* or intramuscularly at day old. Microscopic lesions in the bursa were investigated, 3 and 10 days after challenge. Protection against challenge was determined and the serological response at the age of 14 days old was determined with the VN-test.

1. Average microscopic lesion score in the bursa 3 and 10 days after challenge.

Days post		Group	
challenge	In ovo	Day old	None-vaccinated
3	3.3	0.0	5.0
10	0.2	0.0	5.0

2. Protection after challenge

		Group	
	In ovo	Day old	None-vaccinated
% protection	91.6	100	0

10

15

20

5

3. Serological response against IBDV

		Group		
	In ovo	Day old	None-vaccinated	
VN-titre	$6.4 \pm 1.7$	$6.4 \pm 1.3$	<4.0 ± 0.0	

VN-titre is expressed as log2 of the dilution. Animals with a titre <4.0 log2 are considered

negative

Conclusions

[0096]

30

- 1 The D78/D78/VP5 strain is a highly attenuated IBD-virus
- 2 The virus strain is very mild
- 3 The virus can induce a serological response
- 4 The virus can induce protection
- The virus strain can be applied by intramuscular injection to 1 day old SPF chickens and *in ovo* to 18-days-old embryonated SPF-eggs

[0097] Second animal experiment. The effect of the vaccine is assessed by measurement of the serological response against IBDV and resistance to challenge obtained from administering a challenge virus, 21 days after administering the Gumboro vaccine. The vaccine (10<sup>5.5</sup> TCID<sub>50</sub>/animal of D78/D78/VP5<sup>-</sup>) was applied via the intramuscular route to 14 days old SPF-chickens. Three, 7, 14, and 20 days after vaccination and 3 days after challenge Bursa, spleen, thymus, liver, duodenum, pancreas, ceacal tonsils and harderian gland were investigated for microscopic lesions. Ten days after challenge Bursae were investigated for microscopic lesions. Sera were tested in the VN-test. And mortality was scored after challenge.

4	5

1. Percentage	e mortality after challenge:
	Mortality after challenge
Vaccinated group	0%
Control group	50%

,,,

	2.	Microscop	ic lesions of	the vaccina	ted group befo	re and after c	hallenge:	
Days post	Bursa	Spleen	Thymus	Liver	Duodeum	Pancreas	Ceacal	Harderian
Vaccinat.							Tonsils	Gland
3	0	0	0	0	0	0	0	0
7	0	0	0	0	0	0	0	0

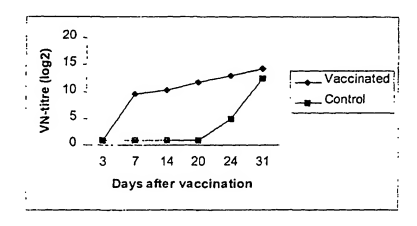
55

#### (continued)

	2.	Microscop	ic lesions of	the vaccina	ated group befo	re and after o	hallenge:	
Days post	Bursa	Spleen	Thymus	Liver	Duodeum	Pancreas	Ceacal	Harderian
Vaccinat.							Tonsils	Gland
14	0	0	0	0	0	0	0	0
20	0	0	0	0	0	0	0	0
24	0,A	0	0	0	0 .	0	0	0
31	0,A	ND	ND	ND	ND	ND	ND	ND

A = None vaccinated animals showed a lymphocytic depletion score of 5.0 (100%) and 4.25, 3 and 10 days after challenge, respectively. ND = not done.

### 3. Serological response after vaccination:



### Conclusions

### [0098]

10

15

20

25

30

35

40

45

- 1. The D78/D78/VP5 strain is a highly attenuated IBD-virus
- 2. The virus strain is very mild and does not induce lesions in organs
- 3. The virus can induce a serological response
- 4. The virus can induce protection

# LEGENDS TO THE FIGURES

[0099] Figure 1 Genomic organization of segment A and segment B of IBDV. The numbers indicate the nucleotide positions of the start, end and coding region on the segments.

[0100] Figure 2 Construction of genomic cDNA clones for the preparation of IBDV/VP5. Plasmid pAD78/EK contains the complete D78 segment A cDNA encoding the polyprotein (VP2-VP4-VP3) and VP5. Plasmid pBP2 contains the complete strain P2 segment B encoding VP1. Mutations were introduced in plasmid pAD78/VP5 altering the methionine start codon for VP5 into arginine and creating an artificial Afl II cleavage site. Recombinant plasmids were linearized with the underlined restriction enzymes, followed by T7 polymerase transcription.

[0101] Figure 3 Construction of genomic cDNA clones for the preparation of IBDV/VP5-2. Plasmid pAD78/EK contains the complete D78 segment A cDNA encoding the polyprotein (VP2-VP4-VP3) and VP5. Plasmid pBD78 contains the complete strain D78 segment B encoding VP1. Mutations were introduced in plasmid pAD78/VP5- altering the methionine start codon for VP5 into glutamic acid and creating an artificial BstBI cleavage site. Further mutations were introduced in the arginine and glutamine codon. Recombinant plasmids were linearized with the underlined restriction

enzymes, followed by T7 polymerase transcription.

[0102] Figure 4 Radioimmunoprecipitation of proteins from CEC infected cells with recombinant IBDV. CEC infected cells with IBDV/VP5 (lanes 1-3), IBDV/EK (lanes 4-6) and uninfected controls were immunoprecipitated with rabbit anti-IBDV serum (lanes 1, 4, 7), rabbit anti-VP5 serum (lanes 2, 5, 8) and mAb DIE 7 (lanes 3, 6, 9). Position of molecular mass markers (M) is indicated. Location of the viral proteins VP2, VP3, VP4 and VP5 are marked.

[0103] Figure 5 Replication kinetics of IBDV/EK and IBDV/VP5-. Infectious titers of supernatants (vertical axis) are determined at the times indicated.

#### SEQUENCE LISTING

[0104]

10

15

20

25

30

35

40

45

50

#### (1) GENERAL INFORMATION:

- (i) APPLICANT:
  - (A) NAME: Azko Nobel N.V.(B) STREET: Velperweg 76
  - (C) CITY: Arnhem
  - (E) COUNTRY: The Netherlands (F) POSTAL CODE (ZIP): 6824 BM (G) TELEPHONE: 0412 666379
  - (H) TELEFAX: 0412 650592
- (ii) TITLE OF INVENTION: Recombinant birnavirus vaccine
  - (iii) NUMBER OF SEQUENCES: 8
  - (iv) COMPUTER READABLE FORM:
    - (A) MEDIUM TYPE: Floppy disk
    - (B) COMPUTER: IBM PC compatible
    - (C) OPERATING SYSTEM: PC-DOS/MS-DOS
    - (D) SOFTWARE: Patent In Release #1.0, Version #1.30 (EPO)
  - (2) INFORMATION FOR SEQ ID NO: 1:
    - (i) SEQUENCE CHARACTERISTICS:
      - (A) LENGTH: 2827 base pairs
      - (B) TYPE: nucleic acid
      - (C) STRANDEDNESS: single
      - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: cDNA
  - (ix) FEATURE:
    - (A) NAME/KEY: CDS (B) LOCATION:112..2745
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 1:

	CCG	CCGC	100		CACG	II A	GIGG	CICC	T CT	ICII	GATG	AT.I	CIGC	CAC	C AT	G AGT	117
															Me	t Ser	
_																1	
5																	
	GAC	ATT	TTC	AAC	AGT	CCA	CAG	GCG	CGA	AGC	ACG	ATC	TCA	GCA	GCG	TTC	165
	Asp	Ile	Phe	Asn	Ser	Pro	Gln	Ala	Arg	Ser	Thr	Ile	Ser	Ala	Ala	Phe	
			5					10					15				
10																	
	GGC	ATA	AAG	CCT	ACT	GCT	GGA	CAA	GAC	GTG	GAA	GAA	CTC	TTG	ATC	CCT	213
													Leu				
		20					25		•			30					
15																	
,,	AAA	GTT	TGG	GTG	CCA	CCT	GAG	GAT	CCG	CTT	GCC	AGC	CCT	AGT	CGA	CTG	261
													Pro				201
	35		-			40		•			45				3	50	
20	GCA	AAG	TTC	CTC	AGA	GAG	AAC	GGC	TAC	AAA	GTT	TTG	CAG	CCA	CGG	тст	309
													Gln				303
		•			55			•	•	60					65		
25	CTG	CCC	GAG	AAT	GAG	GAG	TAT	GAG	ACC	GAC	CAA	ATA	CTC	CCA	GAC	TTA	357
													Leu				,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,
				70			•		75	•				80	<i>E</i>		
30	GCA	TGG	ATG	CGA	CAG	ATA	GAA	GGG	GCT	GTT	TTA	AAA	CCC	ACT	СТА	тст	405
50													Pro				103
		•	85	J				90					95				
	CTC	CCT	ATT	GGA	GAT	CAG	GAG	TAC	TTC	CCA	AAJ	TAC	TAC	CCA	ACA	CAT	453
35													Tyr				.,,
		100		•	•		105	•			•	110					
	CGC	ССТ	AGC	AAG	GAG	AAG	CCC	AAT	GCG	TAC	CCG	CCA	GAC	ATC	GCA	СТА	501
40	_	_	_						_				Asp				
	115			•		120				•	125					130	
	CTC	AAG	CAG	ATG	ATT	TAC	CTG	TTT	CTC	CAG	GTT	CCA	GAG	GCC	AAC	GAG	549
45													Glu				
.0					135	•				140					145		
	GGC	CTA	AAG	GAT	GAA	GTA	ACC	CTC	TTG	ACC	CAA	AAC	ATA	AGG	GAC	AAG	597
													Ile				
50				150					155					160	-	•	
	GCC	TAT	GGA	AGT	GGG	ACC	TAC	ATG	GGA	CAA	GCA	AAT	CGA	CTT	GTG	GCC	645
	Ala	Tyr	Gly	Ser	Gly	Thr	Tyr	Met	Gly	Gln	Ala	Asn	Arg	Leu	Val	Ala	
55			165					170					175				

•	•	ATG	AAG	GAG	GTC	GCC	ACT	GGA	AGA	AAC	CCA	AAC	AAG	GAT	CCT	CTA	AAG	693
			Lys															
			180					185					190	_			-	
5																	)	
		CTT	GGG	TAC	ACT	TTT	GAG	AGC	ATC	GCG	CAG	CTA	CTT	GAC	ATC	ACA	CTA	741
			Gly															
		195					200					205					210	
10																		
		CCG	GTA	GGC	CCA	CCC	GGT	GAG	GAT	GAC	AAG	CCC	TĠG	GTG	CCA	CTC	ACA	789
		Pro	Val	Gly	Pro	Pro	Gly	Glu	Asp	Asp	Lys	Pro	Trp	Val	Pro	Leu	Thr	
						215					220					225		
15																		
,,,		AGA	GTG	CCG	TCA	CGG	ATG	TTG	GTG	CTG	ACG	GGA	GAC	GTA	GAT	GGC	GAC	837
		Arg	Val	Pro	Ser	Arg	Met	Leu	Val	Leu	Thr	Gly	Asp	Val	Asp	Gly	Asp	
					230					235					240	-	-	
20		TTT	GAG	GTT	GAA	GAT	TAC	CTT	CCC	AAA	ATC	AAC	CTC	AAG	TCA	TCA	AGT	885
		Phe	Glu	Val	Glu	Asp	Tyr	Leu	Pro	Lys	Ile	Asn	Leu	Lys	Ser	Ser	Ser	
				245					250	•				255				
25		GGA	CTA	CCA	TAT	GTA	GGT	CGC	ACC	AAA	GGA	GAG	ACA	ATT	GGC	GAG	ATG	933
		Gly	Leu	Pro	Tyr	Val	Gly	Arg	Thr	Lys	Gly	Glu	Thr	Ile	Gly	Glu	Met	
			260					265					270					
30		ATA	GCT	ATC	TCA	AAC	CAG	TTT	CTC	AGA	GAG	CTA	TCA	ACA	CTG	TTG	AAG	981
		Ile	Ala	Ile	Ser	Asn	Gln	Phe	Leu	Arg	Glu	Leu	Ser	Thr	Leu	Leu	Lys	
		275					280					285					290	
25			GGT															1029
35		Gln	Gly	Ala	Gly	Thr	Lys	Gly	Ser	Asn	Lys	Lys	Lys	Leu	Leu	Ser	Met	
						295					300					305		
			AGT									-						1077
40		Leu	Ser	Asp	•	Trp	Tyr	Leu	Ser	_	Gly	Leu	Leu	Phe		Lys	Ala	
					310					315					320			
					~ . ~													
			AGG															1125
45		GIU	Arg		Asp	rys	ser	Inr		ren	Thr	гÀг	Thr		Asn	TTE	Trp	
				325					330					335				
		mar.	C C C T	CC2	<b></b>	202		a										
			GCT Ala															1173
50		ser		PIO	ser	PIO	IIII		теп	Met	TTE	ser		116	Int	Trp	Pro	
			340					345					350					
		GTG	ATG	TCC	חמת	AGC	CCA	דעע	ממ	GTG	ጥጥር	ממ	עיייים	GAA	GGG	ተርታጥ	CCA	1221
		_	Met															1221
55		355					360			-41	Deu	365			y	-ys	370	
55		<del>-</del>															3.0	

	TCA	CTC	TAC	AAA	TTC	AAC	CCG	TTC	AGA	GGA	GGG	TTG	AAC	AGG	ATC	GTC	1269
	Ser	Leu	Tyr	Lys	Phe	Asn	Pro	Phe	Arg	Gly	Gly	Leu	Asn	Arg	Ile	Val	
					375					380					385		
5																	
	GAC	TGG	ATA	TTG	GCC	CCG	GAA	GAA	CCC	AAG	GCT	CTT	GTA	TAT	GCG	GAC	1317
			Ile														
		•		390					395	-, 5		200		400		,,,,,	
									555					+00			
10	AAC	מידמי	TAC	ልጥጥ	GTC	CAC	ጥሮአ	אאכ	N.C.C.	TCC	ጥአር	TON	א מייני	CAC	CT X	CAC	1265
																	1365
	7.51.		Tyr 405		vaı	UIS	Ser		1111	rrp	IYL	261		Asp	Leu	GIU	
			403					410					415				
15	אאכ	CCT	CAC	CCA	2.20	maa	».cm	999	<i>a</i>	010						-1.5	
			GAG														1413
	nys	420	Glu	Ala	Asn	Cys		Arg	GIn	HIS	Met		Ala	Ala	Met	Tyr	
		420					425					430					
20	ma c	2002	ama	200		~~~											
20			CTC														1461
			Leu	inr	Arg		Trp	ser	Asp	Asn		Asp	Pro	Met	Phe		
	435					440					445					450	
	~~ ~																
25			TGG														1509
	Gln	Thr	Trp	Ala		Phe	Ala	Met	Asn	Ile	Ala	Pro	Ala	Leu	Val	Val	
					455					460					465		
30			TCG														1557
00	Asp	Ser	Ser	Cys	Leu	Ile	Met	Asn	Leu	Gln	Ile	Lys	Thr	Tyr	Gly	Gln	
				470					475					480			
	GGC	AGC	GGG	AAT	GCA	GCC	ACG	TTC	ATC	AAC	AAC	CAC	CTC	TTG	AGC	ACA	1605
35	Gly	Ser	Gly	Asn	Ala	Ala	Thr	Phe	Ile	Asn	Asn	His	Leu	Leu	Ser	Thr	
			485					490					495				
	CTA	GTG	CTT	GAC	CAG	TGG	AAC	CTG	ATG	AGA	CAG	CCC	AGA	CCA	GAC	AGC	1653
40	Leu	Val	Leu	Asp	Gln	Trp	Asn	Leu	Met	Arg	Gln	Pro	Arg	Pro	Asp	Ser	
		500					505					510					
	GAG	GAG	TTC	AAA	TCA	TTA	GAG	GAC	AAG	CTA	GGT	ATC	AAC	TTT	AAG	ATT	1701
	Glu	Glu	Phe	Lys	Ser	Ile	Glu	Asp	Lys	Leu	Gly	Ile	Asn	Phe	Lys	Ile	
45	515					520					525					530	•
	GAG	AGG	TCC	ATT	GAT	GAT	ATC	AGG	GGC	AAG	CTG	AGA	CAG	CTT	GTC	CTC	1749
	Glu	Arg	Ser	Ile	Asp	Asp	Ile	Arg	Gly	Lys	Leu	Arg	Gln	Leu	Val	Leu	
50					535					540					545		
•																	
	CTT	GCA	CAA	CCA	GGG	TAC	CTG	AGT	GGG	GGG	GTT	GAA	CCA	GAA	CAA	TCC	1797
			Gln														
55				550	-	-			555	-				560			
55																	

•	AGC	CCA	ACT	GTT	GAG	CTT	GAC	CTA	CTA	GGG	TGG	TCA	GCT	ACA	TAC	AGC	1845
	Ser	Pro	Thr	Val	Glu	Leu	Asp	Leu	Leu	Gly	Trp	Ser	Ala	Thr	Tyr	Ser	
			565					570					575				
5																	
			CTC														1893
	Lys		Leu	Gly	Ile	Tyr		Pro	Val	Leu	Asp		Glu	Arg	Leu	Phe	
		580					585					590					
10	TOT	TOT	CCT	ccc	ጥለጥ	ccc	. , , , ,	CON	OMA	G 2 G		120	) Cm	oma			
			GCT Ala														1941
	595	501	AI a	ATG	TÄT	600	пåр	GIY	Val	GIU	605	гÀг	Ser	Leu	rys		
						000					603					610	
15	AAA	GTC	GGG	ATC	GAG	CAG	GCA	TAC	AAG	GTA	GTC	AGG	TAT	GAG	GCG	ፐፐር	1989
•			Gly														1303
			_		615			•	•	620					625		
20	AGG	TTG	GTA	GGT	GGT	TGG	AAC	TAC	CCA	CTC	CTG	AAC	AAA	GCC	TGC	AAG	2037
	Arg	Leu	Val	Gly	Gly	Trp	Asn	Tyr	Pro	Leu	Leu	Asn	Lys	Ala	Cys	Lys	
				630					635					640			
25			GCA														2085
	ASN	Asn	Ala 645	GIÀ	АТА	ATA	Arg	650	HIS	Leu	GIU	Ala		GIY	Phe	Pro	
			043					650					655				
	CTC	GAC	GAG	TTC	CTA	GCC	GAG	TGG	тст	GAG	CTG	TCA	GAG	TTC	GGT	GAG	2133
30			Glu														
		660					665	-				670			•		
•	GCC	TTC	G AA	GGC	TTC	TAA	ATC	AAG	CTG	ACC	GTA	ACA	TCT	GAG	AGC	CTA	2181
35	Ala	Phe	Glu	Gly	Phe	Asn	Ile	Lys	Leu	Thr	Val	Thr	Ser	Glu	Ser	Leu	
	675					680					685					690	
	000	<b>G</b>	ama			001	O.T. N	~~~	000		~~~	~~~					
			CTG														2229
40	AIA	GIU	Leu	ASII	695	PIO	vaı	PIO	PIO	700	PIO	PIO	ASII	vai	705	Arg	
										, 00					703		
	CCA	GTC	AAC	ACT	GGG	GGA	CTC	AAG	GCA	GTC	AGC	AAC	GCC	CTC	AAG	ACC	2277
.2			Asn														
45				710					715					720			
			TAC														2325
50	Gly	Arg	Tyr	Arg	Asn	Glu	Ala		Leu	Ser	Gly	Leu		Leu	Leu	Ala	
50			725					730					735				
	ACA	GCA	AGA	AGC	ССТ	CTG	440	тар	GCA	<del>ር</del> ምጥ	244	GCC	አልር	GCA	מממ	CCC	7777
			Arg														2373
55		740	5		5		745				-15	750			JIU	.114	
55							_					- •					

	GAG	AAA	CTC	CAC	AAG	TCC	AAG	CCA	GAC	GAC	CCC	GAT	GCA	GAC	TGG	TTC	2421
						Ser											
5	755					760					765	•		•		770	
	G3.3	202															
						CTG											2469
	GIU	Arg	ser	GIU		Leu	Ser	Asp	Leu		Glu	Lys	Ala	Asp		Ala	
10					775					780					785		
	AGC	AAG	GTC	GCC	CAC	TCA	GCA	CTC	GTG	GAA	ACA	AGC	GAC	GCC	CTT	GAA	2517
						Ser											``
15				790					795				•	800			
15																	
						TCC											2565
	Ala	Val		Ser	Thr	Ser	Val		Thr	Pro	Lys	Tyr	Pro	Glu	Val	Lys	
20			805					810					815				
	AAC	CCA	CAG	ACC	GCC	TCC	ממ	CCC	CTT	GTT	GGG	OTTC	CNC	CTC	000	222	0.57.0
						Ser											2613
		82Ó					825				OI1	830		Deu	710	A1a	
25																	
	AAG	AGA	GCC	ACC	GGT	GTC	CAG	GCC	GCT	CTT	CTC	GGA	GCA	GGA	ACG	AGC	2661
	Lys	Arg	Ala	Thr	Gly	Val	Gln	Ala	Ala	Leu	Leu	Gly	Ala	Gly	Thr	Ser	
	835					840					845					850	
30																	
						GAG											2709
	Arg	Pro	Met	Gly		Glu	Ala	Pro	Thr		Ser	Lys	Asn	Ala		Lys	
25					855					860					865		
35	ATG	GCC	AAA	CGG	CGG	CAA	CGC	$C\Delta\Delta$	AAG	GAG	<b>∆</b> GC	CGC	<b>ጥ</b> አ አ ረ	יאפפפ	יאיני		2255
						Gln							IAAC	AGCC	.A1		2755
			•	870	_				875			5					
10																	
	GATO	GGAA	ACC A	CTC	AGA	AG AG	GACA	CTAP	TCC	CAGA	CCC	CGTA	TCCC	CG G	CCTI	CGCCT	2815
	GCGC	GGGC	cc c	C													2827
<b>1</b> 5																	
	(2)	INIEOT		ION E	OD 61	=O ID	NO: 0	_									
	(2)	INCO	TAIVIZ	ION F	0K 21	EQ ID	NO: 2	:									

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 878 amino acids
  - (B) TYPE: amino acid
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein

50

55

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 2:

	1	261	.105	110	5	Wali	261	PLO	GIII	10	AIG	Set	1111	116	15	Ala
5	Ala	Phe	Gly	Ile 20	Lys	Pro	Thr	Ala	Gly 25	Gln	Asp	Val	Glu	Glu 30	Leu	Leu
10	Ile	Pro	Lys 35	Val	Trp	Val	Pro	Pro 40	Glu	Asp	Pro	Leu	Ala 45	Ser	Pro	Ser
15	Arg	Leu 50	Ala	Lys	Phe	Leu	Arg 55	Glu	Asn	Gly	Tyr	Lys 60	Val	Leu	Gln	Pro
	Arg 65	Ser	Leu	Pro	Glu	Asn 70	Glu	Glu	Tyr	Glu	Thr 75	Asp	Gln	Ile	Leu	Pro 80
20	Asp	Leu	Ala	Trp	Met 85	Arg	Gln	Ile	Glu	Gly 90	Ala	Val	Leu	Lys	Pro 95	Thr
25	Leu	Ser	Leu	Pro 100	Ile	Gly	Asp	Gln	Glu 105	Tyr	Phe	Pro	Lys	Tyr 110	Tyr	Pro
	Thr	His	Arg 115	Pro	Ser	Lys	Glu	Lys 120	Pro	Asn	Ala	Tyr	Pro 125	Pro	Asp	Ile
30	Ala	Leu 130	Leu	Lys	Gln	Met	Ile 135	Tyr	Leu	Phe	Leu	Gln 140	Val	Pro	Glu	Ala
35	Asn 145	Glu	Gly	Leu	Lys	Asp 150	Glu	Val	Thr	Leu	Leu 155	Thr	Gln	Asn	Ile	Arg 160
	Asp	Lys	Ala	Tyr	Gly 165	Ser	Gly	Thr	Tyr	Met 170	Gly	Gln	Ala	Asn	Arg 175	Leu
40			Met	180					185	_				190		
45			Leu 195					200					205		_	
50		210	Pro				215					220				
	225		Arg			230					235					240
55	Gly	Asp	Phe	Glu	Val 245	Glu	Asp	Tyr	Leu	Pro 250	Lys	Ile	Asn	Leu	Lys 255	Ser

	ser	ser	GIY	260	Pro	Tyr	vai	GIY	Arg 265	Thr	Lys	GIY	Glu	7nr 270	Ile	Gly
	Glu	Met	Ile 275		Ile	Ser	Asn	Gln 280	Phe	Leu	Arg	Glu	Leu 285	Ser	Thr	Leu
10	Leu	Lys 290	Gln	Gly	Ala	Gly	Thr 295	Lys	Gly	Ser	Asn	Lys 300	Lys	Lys	Leu	Leu
15	Ser 305	Met	Leu	Ser	Asp	Tyr 310	Trp	Tyr	Leu	Ser	Cys 315	Gly	Leu	Leu	Phe	Pro 320
	Lys	Ala	Glu	Arg	Tyr 325	Asp	Lys	Ser	Thr	Trp 330	Leu	Thr	Lys	Thr	Arg 335	Asn
20	Ile	Trp	Ser	Ala 340	Pro	Ser	Pro	Thr	His 345	Leu	Met	Ile	Ser	Met 350	Ile	Thr
25	Trp	Pro	Val 355	Met	Ser	Asn	Ser	Pro 360	Asn	Asn	Val	Leu	Asn 365	Ile	Glu	Gly
	Cys	Pro 370	Ser	Leu	Tyr	Lys	Phe 375	Asn	Pro	Phe	Arg	Gly 380	Gly	Leu	Asn	Arg
30	Ile 385	Val	Glu	Trp	Ile	Leu 390	Ala	Pro	Glu	Glu	Pro 395	Lys	Ala	Leu	Val	Tyr 400
35	Ala	Asp	Asn	Ile	Tyr 405	Ile	Val	His	Ser	Asn 417	Thr	Trp	Tyr	Ser	Ile 415	Asp
	Leu	Glu	Lys	Gly 420	Glu	Ala	Asn	Cys	Thr 425	Arg	Gln	His	Met	Gln 430	Ala	Ala
40	Met	Tyr	Tyr 435	Ile	Leu	Thr	Arg	Gly 440	Trp	Ser	Asp	Asn	Gly 445	Asp	Pro	Met
45		450	Gln				455					460				
50	465		Asp			470					475					480
			Gly		485					490					495	
55	Ser	Thr	Leu	Val 500	Leu	Asp	Gln	Trp	Asn 505	Leu	Met	Arg	Gln	Pro 510	Arg	Pro

	Asp	Ser	515		Pne	. ràs	Ser	520		. Asp	Lys	Leu	525		e Asr	) Phe
5	Lys	1le 530		Arg	Ser	·Ile	8 Asp		) Ile	e Arg	Gly	Lys 540		Arg	g Gln	Leu
10	Val 545		Leu	Ala	Gln	. Pro 550		Tyr	Leu	. Ser	Gly 555	Gly	Val	Glu	Pro	Glu 560
15	Gln	Ser	Ser	Pro	Thr 565		Glu	Leu	Asp	Leu 570	Leu	Gly	Trp	Ser	Ala 575	
	Tyr	Ser	Lys	Asp 580	Leu	Gly	Ile	Tyr	Val 585	Pro	Val	Leu	Asp	Lys 590	Glu	Arg
20	Leu	Phe	Cys 595	Ser	Ala	Ala	Tyr	Pro 600	Lys	Gly	Val	Glu	Asn 605	Lys	Ser	Leu
25	Lys	Ser 610	Lys	Val	Gly	Ile	Glu 615	Gln	Ala	Tyr	Lys	Val 620	Val	Arg	Tyr	Glu
	Ala 625	Leu	Arg	Leu	Val	Gly 630	Gly	Trp	Asn	Tyr	Pro 635	Leu	Leu	Asn	Lys	Ala 640
30	Cys	Lys	Asn	Asn	Ala 645	Gly	Ala	Ala	Arg	Arg 650	His	Leu	Glu	Ala	Lys 655	Gly
35	Phe	Pro	Leu	Asp 660	Glu	Phe	Leu	Ala	Glu 665	Trp	Ser	Glu	Leu	Ser 670	Glu	Phe
	Gly	Glu	Ala 675	Phe	Glu	Gly	Phe	Asn 680	Ile	Lys	Leu	Thr	Val 685	Thr	Ser	Glu
40		Leu 690	Ala	Glu	Leu	Asn	Lys 695	Pro	Val	Pro	Pro	Lys 700	Pro	Pro	Asn	Val
45	Asn 705	Arg	P <b>5</b> 0	Val	Asn	Thr 710	Gly	Gly	Leu	Lys	Ala 715	Val	Ser	Asn	Ala	Leu 720
50	Lys	Thr	Gly		Tyr 725	Arg	Asn	Glu	Ala	Gly 730	Leu	Ser	Gly	Leu	Val 735	Leu
	Leu	Ala		Ala 740	Arg	Ser	Arg	Leu	Gln 745	Asp	Ala	Val	Lys	Ala 750	Lys	Ala
55	Glu	Ala	Glu 755	Lys	Leu	His		Ser 760	Lys	Pro	Asp		Pro 765	Asp	Ala	Asp

	Trp	770	Glu	Arg	Ser	Glu	775	Leu	Ser	Asp	Leu	Leu 780	Glu	Lys	Ala	Asp
5	Ile 785	Ala	Ser	Lys	Val	Ala 790	His	Ser	Ala	Leu	Val 795	Glu	Thr	Ser	Asp	Ala 800
10	Leu	Glu	Ala	Val	Gln 805	Ser	Thr	Ser	Val	Tyr 810		Pro	Lys	Tyr	Pro 815	Glu
15	· Val	Lys	Asn	Pro 820	Gln	Thr	Ala	Ser	Asn 825	Pro	Val	Val	Gly	Leu 830	His	Leu
	Pro	Ala	Lys 835	Arg	Ala	Thr	Gly	Val 840	Gln	Ala	Ala	Leu	Leu 845	Gly	Ala	Gly
20	Thr	ser 850	Arg	Pro	Met	Gly	Met 855	Glu	Ala	Pro	Thr	Arg 860	Ser	Lys	Asn	Ala
25	Val 865	Lys	Met	Ala	Lys	Arg 870	Arg	Gln	Arg	Gln	Lys 875	Glu	Ser	Arg		
30	(2) INFORM															
35	(B) (C)	LENG TYPE: STRA	TH: 32 nucle NDED	261 ba lic acid	se pai l : singl	irs										
40	(ii) MOL (ix) FEA			PE: cD	NA						•					
45	(B) I	LOCA	KEY:	9753		ı. e=0		<b>.</b> . 2.								
	(xi) SEQ	OENC	)E DE	SCRI	TION	: SEQ	אוטווי	J: 3:								
50	GGATACGATC	GGT	CTGAC	ccc c	GGGG	GAGT	C ACC	CCGGG	GAC	AGGC	CGTC	AA GG	CCTT	GTTC		60
55	CAGGATGGGA	CTC	CTCC	TTC T	'ACAA	CGCT	A TC			GTT A						114
-																

	ACA	AAC	GAT	CGC	AGC	GAT	GAC	AAA	CCT	GCA	AGA	TCA	AAC	CCA	ACA	GAT	162
	Thr	Asn	Asp	Arg	Ser	Asp	Asp	Lys	Pro	Ala	Arg	Ser	Asn	Pro	Thr	Asp	
				10					15					20			
5																	
	TGT	TCC	GTT	CAT	ACG	GAG	CCT	TCT	GAT	GCC	AAC	AAC	CGG	ACC	GGC	GTC	210
	Cys	Ser	Val	His	Thr	Glu	Pro	Ser	Asp	Ala	Asn	Asn	Arg	Thr	Gly	Val	
			25					30					35				
10																	
	CAT	TCC	GGA	CGA	CAC	CCT	GGA	GAA	GCA	CAC	TCT	CAG	GTC	AGA	GAC	CTC	258
	His	Ser	Gly	Arg	His	Pro	Gly	Glu	Ala	His	Ser	Gln	Val	Arg	Asp	Leu	
		40	_	_			45					50					
15																	
15	GAC	CTA	CAA	TTT	GAC	TGT	GGG	GGA	CAC	AGG	GTC	AGG	GCT	AAT	TGT	CTT	306
		Leu															
	55				-	60	_	_		_	65	_			-	70	
20	TTT	CCC	TGG	ATT	ccc	TGG	CTC	AAT	TGT	GGG	TGC	TCA	CTA	CAC	ACT	GCA	354
	Phe	Pro	Trp	Ile	Pro	Trp	Leu	Asn	Cys	Gly	Cvs	Ser	Leu	His	Thr	Ala	
			•		75	•			•	80	•				85		
25	GGG	CAA	TGG	GAA	CTA	CAA	GTT	CGA	TCA	GAT	GCT	CCT	GAC	TGC	CCA	GAA	402
	Gly	Gln	Trp	Glu	Leu	Gln	Val	Arg	Ser	Asp	Ala	Pro	Asp	Cys	Pro	Glu	
	•			90				_	95	•			-	100			
30	CCT	ACC	GGC	CAG	TTA	CAA	CTA	CTG	CAG	GCT	AGT	GAG	TCG	GAG	TCT	CAC	450
30	Pro	Thr	Gly	Gln	Leu	Gln	Leu	Leu	Gln	Ala	Ser	Glu	Ser	Glu	Ser	His	
			105					110					115				
	AGT	GAG	GTC	AAG	CAC	ACT	TCC	TGG	TGG	CGT	TTZ.	TGC	ACT	AAA	CGG	CYC .	498
35	Ser	Glu	Val	Lys	His	Thr	Ser	Trp	Trp	Arg	Leu	Cys	Thr	Lys	Arg	His	
		120		•			125	_	·			130					
	CAT	AAA	CGC	CGT	GAC	CTT	CCA	AGG	AAG	CCT	GAG	TGA	CTG	CA C	SATG	TAGCT	551
40	His	Lys	Arg	Arg	Asp	Leu	Pro	Arg	Lys	Pro	Glu						
	135	•	•	_	_	140					145						
	ACA	ATGGG	TT C	ATGI	CTG	CA AC	AGC	CAACA	TCF	ACG	ACAA	AATT	rggg <i>i</i>	AAC (	TCCI	CAGTAG	611
45																	
	GGG	AAGGO	GT C	ACC	TCCI	C AC	CTT	ACCCA	A CAT	CATA	ATGA	TCTT	rggg1	TAT	STGAC	SGCTTG	671
	GTG#	ACCCC	TAC	rccc	CAA	ra go	GCT	rgaco	CA	CAAA	rggt	AGC	CACA	rgt (	GACAC	GCAGTG	731
50	ACAC	GCCC	CAG A	GTCT	CACAC	CC AT	CAAC	rgcac	3 CCC	BATG	ATTA	CCA	ATTCT	CA 1	rcac?	AGTACC	791
	AAC	CAGGT	rgg (	GTA	ACAA!	C AC	CACTO	TTC:	CAC	CCA	ACAT	TGAT	rgcci	ATC I	ACAA	CCTCA	851
55	GCG'	TTGG	agg 1	AGAGO	CTCG	rg T7	TCA	AACA!	A GCC	TCC	ACGG	CCT	CGTAC	CTG (	GCG	CACCA	911

	TETACCICAL	AGCCITIGAT	GGGACAACGG	TAATCACCAG	GGC1G1GGCC	GCAAACAATG	971
5	GGCTGACGAC	CGGCACCGAC	AACCTTATGC	CATTCAATCT	TGTGATTCCA	ACAAACGAGA	1031
	TAACCCAGCC	AATCACATCC	ATCAAACTGG	AGATAGTGAC	CTCCAAAAGT	GGTGGTCAGG	1091
10	CAGGGGATCA	GATGTCATGG	TCGGCAAGAG	GGAGCCTAGC	AGTGACGATC	CATGGTGGCA	1151
	ACTATCCAGG	GGCCCTCCGT	CCCGTCACGC	TAGTGGCCTA	CGAAAGAGTG	GCAACAGGAT	1211
15	CCGTCGTTAC	GGTCGCTGGG	GTGAGCAACT	TCGAGCTGAT	CCCAAATCCT	GAACTAGCAA	1271
	AGAACCTGGT	TACAGAATAC	GGCCGATTTG	ACCCAGGAGC	CATGAACTAC	ACAAAATTGA	1331
20	TACTGAGTGA	GAGGGACCGT	CTTGGCATCA	AGACCGTCTG	GCCAACAAGG	GAGTACACTG	1391
	ACTTTCGTGA	ATACTTCATG	GAGGTGGCCG	ACCTCAACTC	TCCCCTGAAG	ATTGCAGGAG	1451
25	CATTCGGCTT	CAAAGACATA	ATCCGGGCCA	TAAGGAGGAT	AGCTGTGCCG	GTGGTCTCCA	1511
	CATTGTTCCC	ACCTGCCGCT	CCCCTAGCCC	ATGCAATTGG	GGAAGGTGTA	GACTACCTGC	1571
20	TGGGCGATGA	GGCACAGGCT	GCTTCAGGAA	CTGCTCGAGC	CGCGTCAGGA	AAAGCAAGAG	1631
30	CTGCCTCAGG	CCGCATAAGG	CAGCTGACTC	TCGCCGCCGA	CAAGGGGTAC	GAGGTAGTCG	1691
	CGAATCTATT	CCAGGTGCCC	CAGAATCCCG	TAGTCGACGG	GATTCTTGCT	TCACCTGGGG	1751
35	TACTCCGCGG	TGCACACAAC	CTCGACTGCG	TGTTA:GAGA	GGGTGCCACG	CTATTCCCTG	1811
	TGGTTATTAC	GACAGTGGAA	GACGCCATGA	CACCCAAAGC	ATTGAACAGC	AAAATGTTTG	1871
40	CTGTCATTGA	AGGCGTGCGA	GAAGACCTCC	AACCTCCATC	TCAAAGAGGA	TCCTTCATAC	1931
	GAACTCTCTC	TGGACACAGA	GTCTATGGAT	ATGCTCCAGA	TGGGGTACTT	CCACTGGAGA	1991
<b>4</b> 5	CTGGGAGAGA	CTACACCGTT	GTCCCAATAG	ATGATGTCTG	GGACGACAGC	ATTATGCTGT	2051
	CCAAAGATCC	CATACCTCCT	ATTGTGGGAA	ACAGTGGAAA	TCTAGCCATA	GCTTACATGG	2111
50						AATGCTTGTG	2171
						CGACTTGGCC	2231
55						GCAACGTTCA	2291
	I CAAACGTTTT	CCCTCACAAT	CCACGCGACT	GGGACAGGCT	CCCCTACCTC	AACCTACCAT	2351

	ACCTTCCACC	CAATGCAGGA	CGCCAGTACC	ACCTTGCCAT	GGCTGCATCA	GAGTTCAAAG	2411
5	AGACCCCCGA	ACTCGAGAGT	GCCGTCAGAG	CAATGGAAGC	AGCAGCCAAC	GTGGACCCAC	2471
	TATTCCAATC	TGCACTCAGT	GTGTTCATGT	GGCTGGAAGA	GAATGGGATT	GTGACTGACA	2531
0	TGGCCAACTT	CGCACTCAGC	GACCCGAACG	CCCATCGGAT	GCGAAATTTT	CTTGCAAACG	2591
	CACCACAAGC	AGGCAGCAAG	TCGCAAAGGG	CCAAGTACGG	GACAGCAGGC	TACGGAGTGG	2651
5	AGGCTCGGGG	CCCCACACCA	GAGGAAGCAC	AGAGGGAAAA	AGACACACGG	ATCTCAAAGA	2711
	AGATGGAGAC	CATGGGCATC	TACTTTGCAA	CACCAGAATG	GGTAGCACTC	AATGGGCACC	2771
20	GAGGGCCAAG	CCCCGGCCAG	CTAAAGTACT	GGCAGAACAC	ACGAGAAATA	,CCGGACCCAA	2831
	ACGAGGACTA	TCTAGACTAC	GTGCATGCAG	AGAAGAGCCG	GTTGGCATCA	GAAGAACAAA	2891
	TCCTAAGGGC	AGCTACGTCG	ATCTACGGGG	CTCCAGGACA	GGCAGAGCCA	CCCCAAGCTT	2951
?5	TCATAGACGA	AGTTGCCAAA	GTCTATGAAA	TCAACCATGG	ACGTGGCCCA	AACCAAGAAC	3011
	AGATGAAAGA	TCTGCTCTTG	ACTGCGATGG	AGATGAAGCA	TCGCAATCCC	AGGCGGGCTC	3071
80	TACCAAAGCC	CAAGCCAAAA	CCCAATGCTC	CAACACAGAG	ACCCCCTGGT	CGGCTGGGCC	3131
	GCTGGATCAG	GACCGTCTCT	GATGAGGACC	TTGAGTGAGG	CTCCTGGGAG	TCTCCCGACA	3191
35	CCACCCGCGC	AGGTGTGGAC	ACCAATTCGG	CCTTACAACA	TCCCAAATTG	GATCCGTTCG	3251
	CGGGTCCCCT						3261

40

### (2) INFORMATION FOR SEQ ID NO: 4:

(i) SEQUENCE CHARACTERISTICS:

45

(A) LENGTH: 145 amino acids

- (B) TYPE: amino acid
- (D) TOPOLOGY: linear

50

55

- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 4:

Met Val Ser Arg Asp Gln Thr Asn Asp Arg Ser Asp Asp Lys Pro Ala 1 5 10 15

	Arg	Ser	Asn	Pro 20	Thr	Asp	Cys	Ser	Val 25	His	Thr	Glu	Pro	Ser 30	Asp	Ala
5	Asn	Asn	Arg 35	Thr	Gly	Val	His	Ser 40	Gly	Arg	His	Pro	Gly 45	Glu	Ala	His
10	Ser	Gln 50	Val	Arg	Asp	Leu	Asp 55	Leu	Gln	Phe	Asp	Cys 60	Gly	Gly	His	Arg
15	Val 65	Arg	Ala	Asn	Суѕ	Leu 70	Phe	Pro	Trp	Ile	Pro 75	Trp	Leu	Asn	Cys	Gly 80
	Cys	Ser	Leu	His	Thr 85	Ala	Gly	Gln	Trp	Glu 90	Leu	Gln	Val	Arg	Ser 95	Asp
20	Ala	Pro	Asp	Cys 100	Pro	Glu	Pro	Thr	Gly 105	Gln	Leu	Gln	Leu	Leu 110	Gln	Ala
25	Ser	Glu	Ser 115	Glu	Ser	His	Ser	Glu 120	Val	Lys	His	Thr	Ser 125	Trp	Trp	Arg
30	Leu	Cys 130	Thr	Lys	Arg	His	His 135	Lys	Arg	Arg	Asp	Leu 140	Pro	Arg	Lys	Pro
	Glu 145															
35	(2) INFORMA															
<b>40</b>	(B) 1	ENCE ENGT YPE: STRAN	H: 320 nuclei	61 bas c acid	e pair	s										
	(D) 1	TOPOL ECULE														
45	(ix) FEAT	TURE:														
50	(B) L	NAME/ .OCAT	ION:1	3131												
	(xi) SEQ	UENC	E DES	CRIP	TION:	SEQ I	D NO	: 5:								

55

	CAG	GATG(	GGA	CTCC	TCCT	TC T	ACAA	CGCT	A TC	ATTG	ATGG	TTA	GTAG	AGA	TCAG	ACAAAC	120
	GAT	CGCA	GCG	ATG	ACA	AAC	CTG	CAA	GAT	CAA	ACC	CAA	CAG	ATT	GTT	CCG	169
5				Met	Thr	Asn	Leu	Gln	Asp	Gln	Thr	Gln	Gln	Ile	Val	Pro	
				1				`5					10				
	TTC	ATA	CGG	AGC	CTT	CTG	ATG	CCA	ACA	ACC	GGA	CCG	GCG	TCC	ATT	CCG	217
10																Pro	
		15					20					25					
	GAC	GNC	እርር	CTG	GAG	አልር	כאכ	እርጥ	CTC	አርር	ጥሮአ	CNC	ACC	TCC	እሮሮ	ሞእር	265
																Tyr	265
15	30					35					40			501	2112	45	
													GTC				313
20	Asn	Leu	Thr	Val		Asp	Thr	Gly	Ser		Leu	Ile	Val	Phe		Pro	
20					50					55					60		
	GGA	TTC	CCT	GGC	TCA	ATT	GTG	GGT	GCT	CAC	TAC	ACA	CTG	CAG	GGC	AAT	361
	Gly	Phe	Pro	Gly	ser	Ile	Val	Gly	Ala	His	Tyr	Thr	Leu	Gln	Gly	Asn	
25				65					70					75			
	GGG	AAC	TAC	AAG	TTC	GAT	CAG	ATG	CTC	CTG	ACT	GCC	CAG	AAC	CTA	CCG	409
	Gly	Asn	Tyr	Lys	Phe	Asp	Gln	Met	Leu	Leu	Thr	Ala	Gln	Asn	Leu	Pro	
30			80					85					90				
													CTC				457
	AIA	ser 95	ıyr	Asn	Tyr	Cys	Arg	Leu	vaı	Ser	Arg	Ser 105	Leu	Inr	vai	Arg	
35		,,					100					103					
	TCA	AGC	ACA	CTT	CCT	GGT	GGC	GTT	TAT	GCA	CTA	AAC	GGC	ACC	ATA	AAC	505
	Ser	Ser	Thr	Leu	Pro	Gly	Gly	Val	Tyr	Ala	Leu	Asn	Gly	Thr	Ile	Asn	
	110					115					120					125	
40	000	CTC	N C C	TT.C	C	CCA	እሮሮ	CTC	አርም	CAA	OTO	ארא	GAT	CTTT	N.C.C	ma.a	
													Asp				553
					130	<b>U</b> -1				135					140	-1-	
45																	
45													AAA				601
	Asn	Gly	Leu		Ser	Ala	Thr	Ala		Ile	Asn	Asp	Lys		Gly	Asn	
				145					150					155			
50	GTC	CTA	GTA	GGG	GAA	GGG	GTC	ACC	GTC	CTC	AGC	TTA	ccc	ACA	TCA	TAT	649
	Val	Leu	Val	Gly	Glu	Gly	Val	Thr	Val	Leu	Ser	Leu	Pro	Thr	Ser	Tyr	
			160					165					170				
	GAT	CTT	GGG	TAT	GTG	AGG	CTT	GGT	GAC	CCC	ATT	CCC	GCA	ATA	GGG	CTT	697
55													_		_	-	

•	•	Asp	Leu 175	Gly	Туг	Val	Arg	Leu 180	Gly	Asp	Pro	Ile	Pro 185	Ala	Ile	Gly	Leu	
5			CCA Pro															745
10			ACC Thr															793
15			GGT Gly															841
20			AGC Ser															889
25			CTT Leu 255															937
30			GTA Val															985
35			GAC Asp															1033
40			CAG Gln					Ile		Leu								1081
45			GGT Gly															1129
			GTG Val 335															1177
50			CTA Leu															1225
55		GCT	GGG	GTG	AGC	AAC	TTC	GAG	CTG	ATC	CCA	AAT	CCT	GAA	CTA	GCA	AAG	1273

• •	Ala	Gly	Val	Ser	Asn 370	Phe	Glu	Leu	Ile	Pro 375	Asn	Pro	Glu	Leu	Ala 380	Lys	
5		CTG Leu															1321
10	Thr	AAA Lys	Leu 400	Ile	Leu	Ser	Glu	Arg 405	Asp	Arg	Leu	Ġly	Ile 410	Lys	Thr	Val	1369
15	Trp	CCA Pro 415	Thr	Arg	Glu	Tyr	Thr 420	Asp	Phe	Arg	Glu	Tyr 425	Phe	Met	Glu	Val	1417
20		GAC Asp															1465
25		ATA Ile															1513
30		TTC Phe															1561
35		TAC Tyr															1609
40		GCG Ala 495				Ala		Ala			Gly						1657
45		CTC Leu															1705
		CCC Pro															1753
50		CGC Arg		_													1801
55	CTA	TTC	CCT	GTG	GTT	ATT	ACG	ACA	GTG	GAA	GAC	GCC	ATG	ACA	CCC	AAA	1849

•	Leu	Phe	Pro 560		Val	Ile	Thr	Thr 565		Glu	Asp	Ala	Met 570		Pro	Lys	
5			AAC Asn														1897
	Omo	575				<b></b>	580				. 0.	585					
10		Gln	CCT Pro				Arg					Arg					1945
4-5	CAC	AGA	GTC	TAT	GGA			CCA	GAT	GGG			CCA	CTG	GAG		1993
15			Val														
20			GAC Asp														2041
	A TOTT	N TOC	CTC.	625					630					635			
25			CTG Leu 640														2089
			GCC Ala														2137
30		655					660					665					
			GCT Ala			Gly					Cys					Lys	2185
35		AGC	TTT	AGA	AGC	675 ACC	AAG	crc	GCC	ACT	680 GCA	CAC	CGA	CTT	GGC	685 CTT	2233
40			Phe														
40			GCT Ala														2281
45	GCA	ACG	TTC	705 ATC	AAA	CGT	TTC	CCT	710 CAC	ААТ	CCA	CGC	GAC	715 TGG	GAC	AGG	2329
			Phe 720														
50			TAC Tyr														2377
55	TAC		CTT	GCC	ATG	GCT		TCA	GAG	TTC	AAA		ACC	ccc	GAA	СТС	2425

•	. Tyr 750		Leu Ala		Ala 755	Ala	Ser	Glu	Phe	Lys 760	Glu	Thr	Pro	Glu	Leu 765	
5			SCC GTC													2473
10			CCT GCA Ser Ala 785													2521
15		Thr A	AC ATG ASP Met													2569
20			AT TTT		Ala											2617
25			AG TAC ys Tyr	Gly '												2665
30			AG GAA lu Glu													2713
35			CC ATG hr Met 865													2761
40		Gly H	AC CGA is Arg 80													2809
40			AA ATA lu Ile		Asp											2857
45			AG AGC ys Ser	Arg :												2905
50			TC TAC le Tyr													2953
55	ATA	GAC G	AA GTT	GCC .	AAA	GTC	TAT	GAA	ATC	AAC	CAT	GGA	CGT	GGC	CCA	3001

	Ile	Asp	Glu	Val 945	Ala	Lys	Val	Tyr	Glu 950	Ile	Asn	His	Gly	Arg 955	Gly	Pro	)		
5											ACT								3049
10 .											CCC							:	3097
15											GGC Gly 1000	Arg						;	3145
20		TCT Ser				Leu		TGAG	GCTC	CT	GGGAG	TCTC	c ca	ACA	CCAC	С		3	3196
25	ccc		TG T	rggac	CACCA	LA TI	CCGGC	CTTA	CAA	CAT	CCCA	AATT	GGAT	cc c	TTC	GCGG	GT		3256
30		INFOF			OR S													3	261
35		(ii) Me	B) TYI D) TO OLEC	PE: ai POLC	I: 101: mino a DGY: I TYPE: DESC	acid inear : prote	ein	ds SEQ I	D NO	: 6:									
40		•								•									
			Met 1	Thr	Asn	Leu	Gln 5	Asp	Gln	Th	r Glr	Glr 10		e Va	l Pr	o P	he	Ile 15	Arg
45			Ser	Leu	Leu	Met 20	Pro	Thr	Thr	Gl	y Pro		a Sei	r Il	e Pr		sp 30	Asp	Thr
50			Leu	Glu	Lys 35	His	Thr	Leu	Arg	Se:	r Glu O	Th:	c Sei	c Th		r A	sn	Leu	Thr
			Val	Gl <i>y</i> 50	Asp	Thr	Gly	Ser	Gly 55		u Ile	e Val	l Phe		e Pr O	:0 G	ly	Phe	Pro
55			Gly	Ser	Ile	Val	Gly	Ala	His	Ty	r Thr	Le	ı Glı	n Gl	y As	n G	ly	Asn	Tyr

•	·	65					70					75					80
5		Lys	Phe	Asp	Gln	Met . 85		Leu	Thr	Ala	Gln 90		Leu	Pro	Ala	Ser 95	-
		Asn	Tyr	Cys	Arg 100	Leu	Val	Ser	Arg	Ser 105		Thr	Val	Arg	Ser 110	Ser	Thr
10		Leu	Pro	Gly 115		Val	Tyr	Ala	Leu 120	Asn	Gly	Thr	Ile	Asn 125	Ala	Val	Thr
15		Phe	Gln 130	Gly	Ser	Leu	Ser	Glu 135	Leu	Thr	Asp	Val	Ser 140	Tyr	Asn	Gly	Leu
20		Met 145	Ser	Ala	Thr	Ala	Asn 150	Ile	Asn	Asp	Lys	Ile 155	Gly	Asn	Val	Leu	Val 160
				Gly		165					170		•			175	
25				Arg	180					185					190		
30				Ala 195					200					205			
			210	Ala				215					220				
35		225		Ile			230					235					240
40				Gly		245					250				-	255	
				Ala	260					265					270		
45				Ala 275					280					285		_	
50			290	Pro Ser				295					300				
<i>EE</i>		305					310					315					320
55		AIA	GTÅ	Asp	GIN	wec	ser	Lp	ser	Ala	Arg	GΙΆ	Ser	Leu	Ala	Val	Thr

					325					330					335	ı
5.	Ile	His	Gly	Gly 340	Asn	Tyr	Pro	Gly	Ala 345		Arg	Pro	Val	Thr 350		Val
10	Ala	Tyr	Glu 355	Arg	Val	Ala	Thr	Gly 360	Ser	Val	Val	Thr	Val 365		Gly	Val
	Ser	Asn 370	Phe	Glu	Leu	Ile	Pro 375	Asn	Pro	Glu	Leu	Ala 380	Lys	Asn	Leu	Val
15	Thr 385	Glu	Tyr	Gly	Arg	Phe 390	Asp	Pro	Gly	Ala	Met 395	Asn	Tyr	Thr	Lys	Leu 400
20	Ile	Leu	Ser	Glu	Arg 405	Asp	Arg	Leu	Gly	Ile 410	Lys	Thr	Val	Trp	Pro 415	Thr
	Arg	Glu	Tyr	Thr 420	Asp	Phe	Arg	Glu	Tyr 425	Phe	Met	Glu	Val	Ala 430	Asp	Leu
25	Asn	Ser	Pro 435	Leu	Lys	Ile	Ala	Gly 440	Ala	Phe	Gly	Phe	Lys 445	Asp	Ile	Ile
30		Ala 450					455					460				
	465	Ala				470					475				_	480
35		Gly			485					490					495	
40		Lys		500					505					510		
45		Asp	515					520					525			
45		Pro 530					535					540				
50	545	His				550					555					560
<i>EE</i>		Val			565					570					<b>5</b> 75	
55	Ser	Lys	Met	Phe	Ala	Val	Ile	Glu	Gly	Val	Arg	Glu	Asp	Leu	Gln	Pro

				580					585					590		
5	Pro	Ser	Gln 595	Arg	Gly	Ser	Phe	Ile 600	Arg	Thr	Leu	Ser	Gly 605	His	Arg	Val
10	Tyr	Gly 610	Tyr	Ala	Pro	Asp	Gly 615	Val	Leu	Pro	Leu	Glu 620	Thr	Gly	Arg	Asp
	Tyr 625	Thr	Val	Val	Pro	Ile 630	Asp	Asp	Val	Trp	Asp 635	Asp	Ser	Ile	Met	Leu 640
15	Ser	Lys	Asp	Pro	lle 645	Pro	Pro	Ile	Val	Gly 650	Asn	Ser	Gly	Asn	Leu 655	Ala
20	Ile	Ala	Tyr	Met 660	Asp	Val	Phe	Arg	Pro 665	Lys	Val	Pro	Ile	His 670	Val	Ala
	Met	Thr	Gly 675	Ala	Leu	Asn	Ala	Cys 680	Gly	Glu	Ile	Glu	Lys 685	Val	Ser	Phe
25		690					695					Gly 700				
30	Gly 705		-			710					715					720
					725					730		Asp			735	
35	Leu			740	_				745		-			750		
40	Ala		755					760					765			
45		<b>7</b> 70					775				_	780				
	Ala 785					790					795					800
50					805					810		His Ser			815	
				820	nia				825	501	~ <b>,</b> 3			830	a	-19
55	Tyr	Gly	Thr	Ala	Gly	Tyr	Gly	Val	Glu	Ala	Arg	Gly	Pro	Thr	Pro	Glu

•	•			835					840					845			
5		Glu	Ala 850	Gln	Arg	Glu	Lys	Asp 855	Thr	Arg	Ile	Ser	Lys 860	Lys	Met	Glu	Thr
10		Met 865	Gly	Ile	Tyr	Phe	Ala 870	Thr	Pro	Glu	Trp	Val 875	Ala	Leu	Asn	Gly	His 880
		Arg	Gly	Pro	Ser	Pro 885	Gly	Gln	Leu	Lys	Tyr 890	Trp	Gln	Asn	Thr	Arg 895	Glu
15		Ile	Pro	Asp	Pro 900	Asn	Glu	Asp	Tyr	Leu 905	Asp	Tyr	Val	His	Ala 910	Glu	Lys
20		Ser	Arg	Leu 915	Ala	Ser	Glu	Glu	Gln 920	Ile	Leu	Arg	Ala	Ala 925	Thr	Ser	Ile
25		Tyr	Gly 930	Ala	Pro	Gly	Gln	Ala 935	Glu	Pro	Pro	Gln	Ala 940	Phe	Ile	Asp	Glu
25		Val 945	Ala	Lys	Val	Tyr	Glu 950	Ile	Asn	His	Gly	Arg 955	Gly	Pro	Asn	Gln	Glu 960
30	•	Gln	Met	Lys	Asp	Leu 965	Leu	Leu	Thr	Ala	Met 970	Glu	Met	Lys	His	<b>Arg</b> 975	Asn
35	:	Pro	Arg	Arg	Ala 980	Leu	Pro	Lys	Pro	Lys 985	Pro	Lys	Pro	Asn	Ala 990	Pro	Thr
	(	Gln	Arg	Pro 995	Pro	Gly	Arg	Leu	Gly 1000	_	Trp	Ile	Arg	Thr 1005		Ser	Asp
40	(	Glu	Asp 1010	Leu )	Glu												
45	(2) INF	ORM	1ATIO	N FOF	R SEQ	ID NO	D: 7:										
	(i)	SEQ	UENC	E CH	ARAC	TERIS	STICS	:									
50		(A) (B)	LENG TYPE	STH: 3 :: nucl	261 b	ase pa	airs										
		(D)	TOPO	DLOG'	Y: line	ar											
55	(ii)	) MOL	.ECUL	E TY	PE: cE	NA											

(ix) FEATURE:

•	. (A) NAME/KEY: CDS (B) LOCATION:97531														
5	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 7:														
	GGATACGATC GGTCTGACCC CGGGGGAGTC ACCCGGGGAC AGGCCGTCAA GGCCTTGTTC	60													
10	CAGGATGGGA CTCCTCCTTC TACAACGCTA TCATTC GAA GTT AGT TGA GAT CTG  Glu Val Ser * Asp Leu  1 5	114													
15	ACA AAC GAT CGC AGC GAT GAC AAA CCT GCA AGA TCA AAC CCA ACA GAT Thr Asn Asp Arg Ser Asp Asp Lys Pro Ala Arg Ser Asn Pro Thr Asp  10 15 20	162													
20	(2) INFORMATION FOR SEQ ID NO: 8:														
	(i) SEQUENCE CHARACTERISTICS:														
25	(A) LENGTH: 2827 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear														
3 <i>0</i>	(ii) MOLECULE TYPE: cDNA														
	(ix) FEATURE:														
35	(A) NAME/KEY: CDS (B) LOCATION:1122745														
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 8:														
10	GGATACGATG GGTCTGACCC TCTGGGAGTC ACGAATTAAC GTGGCTACTA GGGGCGATAC	60													
	CCGCCGCTGG CTGCCACGTT AGTGGCTCCT CTTCTTGATG ATTCTGCCAC C ATG AGT Met Ser	117													
<b>1</b> 5	1														
50 .	GAC ATT TTC AAC AGT CCA CAG GCG CGA AGC ACG ATC TCA GCA GCG TTC Asp Ile Phe Asn Ser Pro Gln Ala Arg Ser Thr Ile Ser Ala Ala Phe 5 10 15	165													
	GGC ATA AAG CCT ACT GCT GGA CAA GAC GTG GAA GAA CTC TTG ATC CCT	213													

		20					25					2.0					
												30					
5							,			•							
3					CCA												261
	Lys	Val	Trp	Val	Pro	Pro	Glu	Asp	Pro	Leu	Ala	Ser	Pro	Ser	Arg	Leu	
	35					40					45					50	
40	CON	220	mma														
10					AGA												309
	Ald	гуs	Pne	rea	Arg 55	GIU	Asn	GIY	Tyr		Vai	Leu	Gln	Pro		Ser	
					33					60					65		
15	CTG	CCC	GAG	AAT	GAG	GAG	TAT	GAG	ACC	GAC	CAA	ATA	CTC	CCA	GAC	ТТА	357
7.5					Glu												•••
				70					75	_				80	-		
20					CAG												405
20	Ala	Trp		Arg	Gln	Ile	Glu		Ala	Val	Leu	Lys		Thr	Leu	Ser	
			85					90					95				
	CTC	CCT	ATT	GGA	GAT	CAG	GAG	ראכי	ጥጥር	רכא	ממ	тас	ጥልሮ	CCA	מכים	CAT	453
25					Asp												453
20		100		•	-		105	-2-			-7-	110	-1-				
																	•
	CGC	CCT	AGC	AAG	GAG	AAG	CCC	AAT	GCG	TAC	CCG	CCA	GAC	ATC	GCA	CTA	501
30	Arg	Pro	Ser	Lys	Glu	Lys	Pro	Asn	Ala	Tyr	Pro	Pro	Asp	Ile	Ala	Leu	
50	115					120					125					130	
					ATT												549
35	rea	rys	GIN	Met	Ile	lyr	ren	Phe	Leu		Va.	Pro	Glu	Ala		Glu	
30					135					140					145		
	GGC	CTA	AAG	GAT	GAA	GTA	ACC	CTC	TTG	ACC	CAA	AAC	ATA	AGG	GAC	AAG	597
					Glu												33,
40				150					155					160	•	•	
					GGG												645
	Ala	Tyr		Ser	Gly	Thr	Tyr	Met	Gly	Gln	Ala	Thr	Arg	Leu	Val	Ala	
45			165					170					175				
40	N TO CO	220	C N C	CMC	000	3 CM	003										
					GCC												693
		180	014	VAI	Ala	TILL	185	Arg	WPII	PIO	ASII	190	ASD	PLO	теп	гÀг	
50							103					170					
	CTT	GGG	TAC	ACT	TTT	GAG	AGC	ATC	GCG	CAG	CTA	CTT	GAC	ATC	ACA	CTA	741
					Phe												
	195					200					205		-			210	
55																	
JJ	CCG	GTA	GGC	CCA	CCC	GGT	GAG	GAT	GAC	AAG	CCC	TGG	GTG	CCA	CTC	ACA	789

•	•	Pro	Val	Gly	Pro	Pro 215	Gly	Glu	Asp	Asp	Lys 220	Pro	Trp	Val	Pro	Leu 225	Thr	
						CGG Arg												837
10						GAT Asp												885
15						GTA Val												933
20						AAC Asn												981
25						ACA Thr 295												1029
30						TGG Trp												1077
35						AAA Lys												1125
40						CCA Pro												1173
45						AGC Ser												1221
٠٠						TTC Phe 375												1269
<b>50</b> .						GCC Ala												1317
55		AAC	ATA	TAC	ATT	GTC	CAC	TCA	AAC	ACG	TGG	TAC	TCA	ATT	GAC	CTA	GAG	1365

•	•	Asn	Ile	Tyr 405	Ile	Val	His	Ser	Asn 410	Thr	Trp	Tyr	Ser	Ile 415	Asp	Leu	Glu	
5	٠	AAG	GGT	GAG	GCA	AAC	TGC	ACT	CGC	.CAA	CAC	ATG	CAA	GCC	GCA	ATG	TAC	1413
		Lys	Gly	Glu	Ala	Asn	Cys	Thr	Arg	Gln	His	Met	Gln	Ala	Ala	Met	Tyr	
			420					425					430					
10					ACC												· <del>-</del>	1461
		_	Ile	Leu	Thr	Arg	_	Trp	Ser	Asp	Asn	•	Asp	Pro	Met	Phe		
		435					440					445					450	
15					GCC													1509
		Gin	Thr	Trp	Ala		Phe	Ala	Met	Asn		Ala	Pro	Ala	Leu		Val	
						455					460					465		
					TGC					_				-				1557
20		Asp	Ser	Ser	Cys	Leu	Ile	Met	Asn		Gln	Ile	Lys	Thr	-	Gly	Gln	
					470					475					480			
					TAA													1605
25		Gly	Ser	_	Asn	Ala	Ala	Thr		Ile	Asn	Asn	His		Leu	Ser	Thr	
				485					490					495				
					GAC													1653
30		Leu	Va1	Leu	Asp	Gin	Trp	Asn 505	ren	met	Arg	GII	510	Arg	Pro	Asp	ser	
			300					<b>J</b> 03					310				•	
					AAA							_						1701
			Glu	?he	Lys	Ser		Glu	Asp	Lys	Leu	_	Ile	Asn	Phe	Lys		
35		515					520					525			•		530	
					ATT					_				_				1749
		Glu	Arg	Ser	Ile		Asp	IIE	Arg	GIY		Leu	Arg	GIN	Leu		Leu	
40						535					540					545		
		CTT	GCA	CAA	CCA	GGG	TAC	CTG	AGT	GGG	GGG	GTT	GAA	CCA	GAA	CAA	TCC	1797
		Leu	Ala	Gln	Pro	Gly	Tyr	Leu	Ser	Gly	Gly	Val	Glu	Pro	Glu	Gln	Ser	
					550	_	-			555	_				560			
45																		
		AGC	CCA	ACT	GTT	GAG	CTT	GAC	CTA	CTA	GGG	TGG	TCA	GCT	ACA	TAC	AGC	1845
		Ser	Pro	Thr	Val	Glu	Leu	Asp		Leu	Gly	Trp	Ser	Ala	Thr	Tyr	Ser	
				565					570					575				
50		מממ	G ກ ጥ	СТС	GGG	እ ጥር	ጥኦጥ	ama	CCG	GTG	ستب	GNC	270	ת תם	CCC	CITI N	diam.	1003
					Gly													1893
		-13	580		1		-1-	585				F	590	-Lu	9	بت لا	* 410	
55		TGT	TCT	GCT	GCG	TAT	CCC	AAG	GGA	GTA	GAG	AAC	AAG	AGT	CTC	AAG	TCC	1941

•	•	Cys 595		Ala	. Ala	Tyr	Pro 600		Gly	Val	Glu	Asn 605		Ser	Leu	Lys	Ser 610	
5							CAG											1989
		Lys	Val	. Gly	Ile	615		Ala	Tyr	Lys	Val 620		Arg	Tyr	Glu	Ala 625	Leu	
10							TGG											2037
		Arg	Leu	Val	Gly 630		Trp	Asn	Tyr	Pro 635		Leu ·	Asn	Lys	Ala 640	Cys	Lys	
15							GCT											2085
		ASN	Asn	A1a 645	GIÀ	Ala	Ala	Arg	Arg 650	His	Leu	Glu	Ala	Lys 655	Gly	Phe	Pro	
							GCC											2133
20		Leu	660	GIU	Pne	Leu	Ala	665	Trp	Ser	Glu	Leu	Ser 670	Glu	Phe :	Gly	Glu	
							AAT											2181
25		675	Pne	GIu	GIÀ	Phe	Asn 680	Ile	rys	Leu	Thr	Val 685	Thr	Ser	Glu	Ser	Leu 690	
							CCA											2229
30		Ala	Glu	Leu	Asn	Lys 695	Pro	Val	Pro	Pro	Lys 700	Pro	Pro	Asn	Val	Asn 705	Arg	
							GGA											2277
35		Pro	Val	Asn	710	GIA	Gly	Leu	Lys	715	Val	Ser	Asn	Ala	Leu 720	Lys	Thr	
		GGT	CGG	TAC	AGG	AAC	GAA	GCC	GGA	CTG	AGT	GGT	CTC	GTÇ	CTT	CTA	GCC	2325
		Gly	Arg		Arg	Asn	Glu	Ala			Ser	Gly	Leu		Leu	Leu	Ala	
40				725		•			730					735				
							CTG											2373
		Thr	Ala 740	Arg	Ser	Arg	Leu	Gln 745	Asp	Ala	Val	Lys	Ala 750	Lys	Ala	Glu	Ala	
45		GAG	AAA	CTC	CAC	AAG	TCC	AAG	CCA	GAC	GAC	CCC	GAT	GCA	GAC	TGG	TTC	2421
			Lys	Leu	His	Lys	Ser	Lys	Pro	Asp	Asp		Asp	Ala	Asp	Trp	Phe	
		755					760					765					770	
50		GAA	AGA	TCA	GAA	ACT	CTG	TCA	GAC	CTT	CTG	GAG	AAA	GCC	GAC	ATC	GCC	2469
		Glu	Arg	Ser	Glu	Thr 775	Leu	Ser	Asp	Leu		Glu	Lys	Ala	Asp		Ala	
						//3					780					785		
55		AGC	AAG	GTC	GCC	CAC	TCA	GCA	CTC	GTG	GAA	ACA	AGC	GAC	GCC	CTT	GAA	2517

	Ser	Lys	Val	Ala 790	His	Ser	Ala	Leu	Val 795	G1u	Thr	Ser	Asp	Ala 800	Leu	Glu		
5								TAC Tyr 810									:	2565
10			CAG					CCC Pro					CAC				:	2613
15								GCC Ala									:	2661
20								CCA Pro									2	2709
25								CAA Gln					TAAC	AGCO	AT		2	2755
30	GATO	GGAA	ACC A	CTCA	AGAA	G AC	GACA	CTAA	TCC	CAGA	.ccc	CGTA	TCCC	CG G	CCTI	CGCCT	2	2815
	GCGG	GGGC	cc c	C.													2	2827

#### Claims

35

- 1. A birnavirus mutant which is not able to produce a native VP5 protein as a result of a mutation in the VP5 gene of the birnavirus genome, **characterised in that** the mutation comprises:
  - (i) a substitution of at least two nucleotides of the start codon of the VP5 gene, and
  - (ii) a stop codon in each of the three open reading frames in the 5'-end of the VP5 gene.
  - 2. A birnavirus mutant according to claim 2, characterised in that the birnavirus is infectious bursal disease virus (IBDV).
- 3. A birnavirus mutant according to claims 1-2, **characterised in that** the mutation is in the genome of a virulent field virus.
  - 4. A birnavirus mutant according to claim 2, **characterised in that** the mutation is in the genome of a vaccine strain, preferably in vaccine strain D78.
- 55 5. A birnavirus mutant according to claims 2-4, **characterised in that** the mutant has a mutated start codon and three stop codons in the 5'-end of the VP5 gene as shown in SEQ ID No: 7.
  - 6. A birnavirus mutant according to claims 2-5, characterised in that the IBDV expresses a chimeric VP2 protein

- comprising virus neutralising epitopes of different antigenic IBDV types.
- 7. A vaccine against a birnavirus infection in animals, **characterised in that** it comprises a birnavirus mutant according to claims 1-6 and a pharmaceutically acceptable carrier.
- 8. A method for the attenuation of virulence of a birnavirus in an animal, comprising the step of introducing a mutation in the VP5 gene as a result of which the birnavirus is not able to produce a VP5 protein.
- 9. A method according to claim 9, wherein the mutation is a substitution.
- 10. A method according to claims 8-9, wherein the birnavirus is infectious bursal disease virus (IBDV).
- 11. A method according to claims 8-10, wherein the mutation is in the genome of a virulent field virus.
- 15 12. A method according to claims 8-11, wherein the mutation comprises a substitution of at least two nucleotides of the start codon of the VP5 gene.
  - 13. A method according to claim 12 wherein the mutation comprises additionally one or more stop codons in the 5'end of the VP5 gene.
  - 14. A method according to claim 13, wherein the mutation comprises a stop codon in each of the three open reading frames.
- 15. A method according to claim 14, wherein the mutation is in the start codon and comprises three stop codons in the 5'-end of the VP5 gene as shown in SEQ ID No: 7.

### Patentansprüche

5

10

20

35

40

50

- Eine Birnavirus Mutante, die aufgrund einer Mutation im VP5 Gen des Birnavirus Genoms kein natives VP5 Protein produzieren kann, dadurch gekennzeichnet, dass die Mutation:
  - (i) eine Substitution von mindestens zwei Nukleotiden des Startcodons des VP5 Gens, und
  - (ii) ein Stopcodon in jedem der drei offenen Leserahmen am 5'-Ende des VP5 Gens.

umfasst.

- Eine Birnavirus Mutante gemäss Anspruch 1, dadurch gekennzeichnet dass das Birnavirus ein Infektiöses Bursitis Virus (IBDV) darstellt.
  - 3. Eine Birnavirus Mutante gemäss Ansprüchen 1-2, dadurch gekennzeichnet, dass die Mutation im Genom eines virulenten Feldvirus ist.
- Eine Birnavirus Mutante gemäss Anspruch 2, dadurch gekennzeichnet, dass die Mutation im Genom eines Impfstoff-Stammes, vorzugsweise im Impfstoff-Stamm D78, ist.
  - 5. Eine Birnavirus Mutante gemäss Ansprüchen 2-4, **dadurch gekennzeichnet, dass** die Mutante ein mutiertes Startcodon und drei Stopcodons am 5'-Ende des VP5-Gens wie in SEQ ID No:7 dargestellt besitzt.
  - 6. Eine Birnavirus Mutante gemäss Ansprüchen 2-5, dadurch gekennzeichnet, dass das IBDV ein chimäres VP2 Protein exprimiert, das Virus-neutralisierende Epitope von unterschiedlichen, antigenen IBDV-Typen umfasst.
- Ein Impfstoff gegen eine Birnavirus Infektion in Tieren, dadurch gekennzeichnet, dass er eine Birnavirus Mutante gemäss Ansprüchen 1-6 und einen pharmazeutisch verträglichen Träger umfasst.
  - Verfahren zur Abschwächung der Virulenz eines Birnavirus in einem Tier, welches den Schritt des Einführens einer Mutation in das VP5 Gen umfasst, als Folge dessen das Birnavirus kein VP5 Protein produzieren kann.

- 9. Verfahren gemäss Anspruch 8, worin die Mutation eine Substitution darstellt.
- 10. Verfahren gemäss Ansprüchen 8-9, worin das Birnavirus ein Infektiöses Bursitis Virus (IBDV) darstellt.
- <sup>5</sup> 11. Verfahren gemäss Ansprüchen 8-10, worin die Mutation im Genom eines virulenten Feldvirus ist.
  - 12. Verfahren gemäss Ansprüchen 8-11, worin die Mutation eine Substitution von mindestens zwei Nukleotiden des Startcodons des VP5 Gens umfasst.
- 13. Verfahren gemäss Anspruch 12, worin die Mutation zusätzlich ein oder mehrere Stopcodons am 5'-Ende des VP5-Gens umfasst.
  - 14. Verfahren gemäss Anspruch 13, worin die Mutation in jedem der drei offenen Leserahmen ein Stopcodon umfasst.
- 15. Verfahren gemäss Anspruch 14, worin die Mutation im Startcodon ist und drei Stopcodons am 5'-Ende des VP5-Gens wie in SEQ ID No:7 dargestellt umfasst.

### Revendications

20

25

- 1. Un mutant du birnavirus qui est incapable de produire une protéine VP5 native, résultant d'une mutation dans le gène VP5 du génome du birnavirus, **caractérisé en ce que** la mutation comprend :
  - i) une substitution d'au moins deux nucléotides du codon de départ du gène VP5, et
  - ii) un codon d'arrêt dans chacun des trois cadres ouverts de lecture dans l'extrémité 5' du gène VP5.
- 2. Un mutant du birnavirus selon la revendication 1, caractérisé en ce que le birnavirus est le virus de la bursite infectieuse aviaire (IBDV: *Infections Bursal Diseuse Virus*).
- 30 3. Un mutant du birnavirus selon la revendication 1 ou 2, caractérisé en ce que la mutation est dans le génome d'un virus sauvage virulent.
  - 4. Un mutant du birnavirus selon la revendication 2, caractérisé en ce que la mutation est dans le génome d'une souche de vaccin, de préférence dans la souche de vaccin D78.

35

- 5. Un mutant du birnavirus selon l'une des revendications 2 à 4, caractérisé en ce que le mutant a un codon de départ ayant subi une mutation et les trois codons d'arrêt dans l'extrémité 5' du gène VP5 tel que représenté dans la SEQ ID n° 7.
- 6. Un mutant du birnavirus selon l'une des revendications 2 à 5, caractérisé en ce que l'IBDV exprime une protéine chimère VP2 comprenant des épitopes neutralisant des virus de différents types antigènes d'IBDV.
  - 7. Un vaccin contre une infection par le birnavirus chez des animaux, caractérisé en ce qu'il comprend un mutant du birnavirus selon l'une des revendications 1 à 6 et un support pharmaceutiquement acceptable.

45

- 8. Un procédé pour atténuer la virulence d'un birnavirus chez un animal, comprenant l'étape consistant à introduire une mutation dans le gène VP5 pour que le birnavirus soit incapable de produire une protéine VP5.
- 9. Un procédé selon la revendication 9, dans lequel la mutation est une substitution.

50

- 10. Un procédé selon la revendication 8 ou 9, dans lequel le birnavirus est le virus de la bursite infectieuse aviaire (IBDV).
- 11. Un procédé selon l'une des revendications 8 à 10, dans lequel la mutation est dans le génome d'un virus sauvage virulent.
- 12. Un procédé selon l'une des revendications 8 à 11, dans lequel la mutation comprend une substitution d'au moins deux nucléotides du codon de départ du gène VP5.

13. Un procédé selon la revendication 12, dans lequel la mutation comprend en outre un ou plusieurs codons d'arrêt dans l'extrémité 5' du gène VP5. 14. Un procédé selon la revendication 13, dans lequel la mutation comprend un codon d'arrêt dans chacun des trois cadres ouverts de lecture. 15. Un procédé selon la revendication 14, dans lequel la mutation est située dans le codon de départ et comprend trois codons d'arrêt dans l'extrémité 5' du gène VP5 tel que représenté dans la SEQ ID nº 7. 

Figure 1

nt 3261 3. -NCR nt 2827 nt 3166 nt 2745 VP2 - VP4 - VP3 VP1 P2 segment B D78 segment A nt 112 nt 131 nt 97 nt 1

Genomic organization of segment A of strain D78 and segment B of strain P2

Figure 2

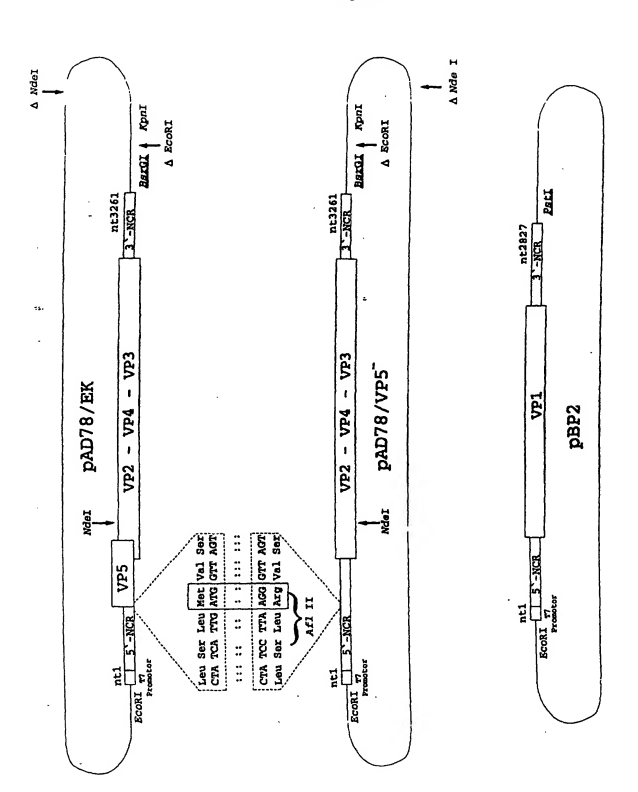


Figure 3

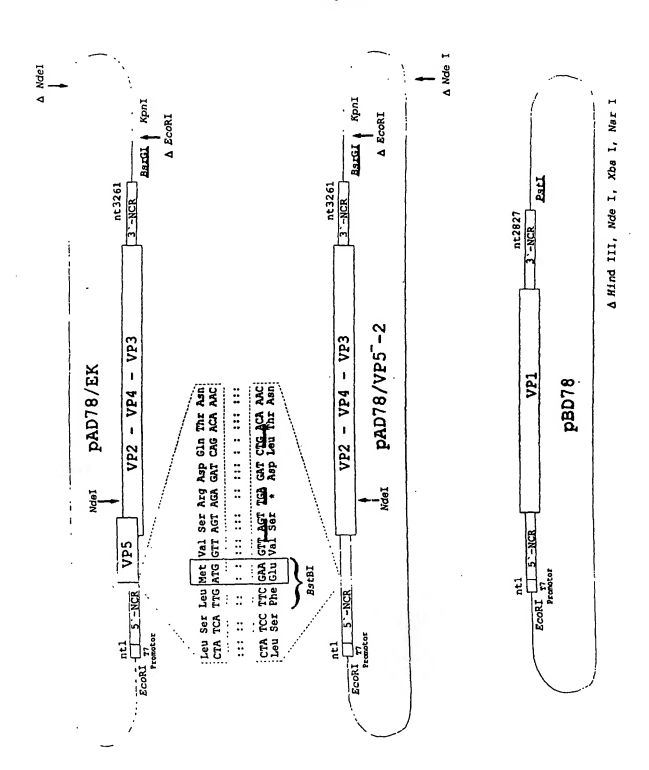


Figure 4

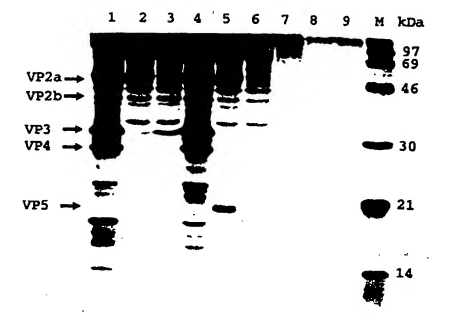
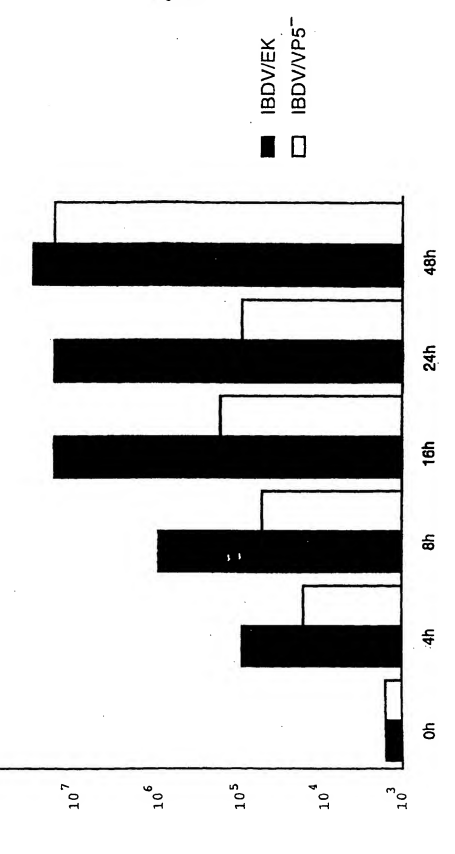


Figure 5



gh